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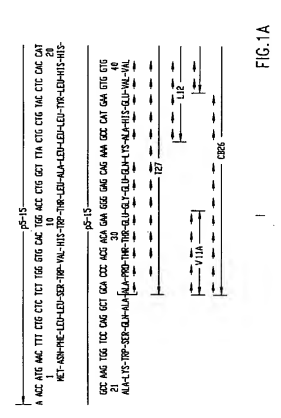
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(1) Applicant: MERCK & CO. INC. 126, East Lincoln Avenue P.O. Box 2000 Rahway New Jersey 07065-0900 (US) 131 Tudor Oval
Westfield, NJ 07090 (US)
Inventor: Thomas, Kenneth A., Jr.
245 Washington Avenue
Chatham Burough, NJ 07928 (US)

(74) Representative: Thompson, John Dr. et al Merck & Co., Inc. European Patent Department Terlings Park Eastwick Road Harlow, Essex CM20 2QR (GB)

- 54) Vascular endothellal cell growth factor C subunit.
- (57) Vascular endothelial cell growth factor C subunit DNA is prepared by polymerase chain reaction techniques. The DNA encodes a protein
 that may exist as either a heterodimer or
 homodimer. The protein is a mammalian vascular endothelial cell mitogen and as such is
 useful for the promotion of vascular development and repair. This unique growth factor is
 also useful In the promotion of tissue repair.



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BRIEF DESCRIPTION OF THE DRAWING

Figure 1. Full length amino acid residue protein translation product and its cDNA coding sequence for VEGF AA subunit A plus polypeptide cleavage products used to determine the amino acid sequence.

Figure 2. Full length amino acid residue protein translation product and its cDNA coding sequence for VEGF AB subunit A plus polypeptide cleavage products used to determine the amino acid sequence.

Figure 3. Full length amino acid residue protein translation product and its cDNA coding sequence for VEGF AB subunit B plus polypeptide cleavage products used to determine the amino acid sequence.

Figure 4. Full cDNA coding sequence and full length amino acid residue protein translation product for VEGF A 146 amino acid residue subunit SEQ ID NOS:23 & 33.

Figure 5. Full cDNA coding sequence and full length amino acid residue protein translation product for VEGF A 190 amino acid residue subunit SEQ ID NOS:30 & 31.

Figure 6. Full cDNA coding sequence and full length amino acid residue protein translation product for VEGF A 214 amino acid residue subunit SEQ ID NOS:34 & 35.

Figure 7. Full cDNA coding sequence and full length amino acid residue protein translation product for VEGF B 138 amino acid residue subunit SEQ ID NOS:36 & 37.

Figure 8. Full cDNA coding sequence and full length amino acid residue protein translation product for VEGF B 158 amino acid residue subunit SEQ ID NOS:38 & 39.

Figure 9. Full cDNA coding sequence and full length amino acid residue protein translation product for VEGF C 154 amino acid residue subunit SEQ ID NOS:40 & 41.

BACKGROUND OF THE INVENTION

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A new class of cell-derived dimeric mitogens with apparently restricted specificity for vascular endothelial cells has recently been identified and generally designated vascular endothelial growth factors (VEGFs). The mitogen has been purified from: conditioned growth media of rat glioma cells, [Conn et al., Proc. Natl. Acad. Sci. USA 87: 1323-1327 (1990)]; conditioned growth media of bovine pituitary folliculo stellate cells [Ferrara and Henzel, Biochem. Biophys. Res. Comm. 161: 851-858 (1989) and Gospodarowicz et al., Proc. Natl. Acad. Sci. USA 86: 7311-7315 (1989)]. An endothelial cell growth factor isolated form mouse neuroblastoma cell line NB41 with an unreduced molecular mass of 43-51 kDa and a reduced mass of 23-29 kDa has been described by Levy et al., Growth Factors 2: 9-19 (1989). Connolly et al. (J. Biol. Chem. 264: 20017-20024 [1989]; J. Clin. Invest, 84: 1470-1478 (1989)) describe a human vascular permeability factor that stimulates vascular endothelial cells to divide in vitro and promotes the growth of new blood vessels when administered into healing rabbit bone grafts or rat corneas. An endothelial cell growth factor has been purified from the conditioned medium of the AtT-20 pituitary cell line by Plouet et al., EMBO Jownal 8: 3801-3806 (1989). The growth factor was characterized as a heterodimer composed of subunits with molecular mass of 23 kDa. Leung et al. (Science 246: 1306-1309 [1989]), Keck et al. (Science 246: 1309-1312 [1989]) and Conn et al. (Proc. Natl. Acad. Sci USA 87: 2628-2632 [1990]) have described cDNAs which encode VEGF A which is homologous to the A and B chains of platelet-derived growth factor. Vascular endothelial growth factor I (VEGF I, VEGF AA) is a homodimer with an apparent molecular mass of 46 kDa, with each subunit having an apparent molecular mass of 23 kDa. VEGF I has distinct structural similarities to platelet-derived growth factor (PDGF), a mitogen for connective tissue cells but not vascular endothelial cells from large vessels.

OBJECTS OF THE INVENTION

It is, accordingly, an object of the present invention to provide novel vascular endothelial growth factor C subunit DNA free of other mammalian DNA. Another object is to provide recombinant genes capable of expressing VEGF C subunit monomer or dimer. Another object is to provide vectors containing the DNA sequences for VEGF A or B plus C subunits. A further object is to provide a host cell transformed with a vector containing the DNA sequence for VEGF A or B plus C or VEGF C alone. It is also an object to provide a recombinant process for making VEGF C subunit. Another object is to provide a novel vascular endothelial cell growth factor which contains the C subunit. This may include heterodimers AC and BC and homodimer CC.

SUMMARY OF THE INVENTION

Vascular endothelial cell growth factor C subunit DNA is prepared by polymerase chain reaction techniques. The DNA encodes a protein that may exist as either a heterodimer or homodimer. The protein is a mammalian vascular endothelial cell mitogen and as such is useful for the promotion of vascular development and

repair. This unique growth factor is also useful in the promotion of tissue repair.

DETAILED DESCRIPTION

The present Invention relates to a unique vascular endothelial cell growth factor (designated VEGF), isolated and purified from glioma cell conditioned medium, which exhibits mitogenic stimulation of vascular endothelial cells. Glioma is defined herein as any neoplasm derived from one of the various types of cells that form the interstitial tissue of the central nervous system including brain, spinal cord, posterior pituitary gland and retina. Consequently, the scope of the present invention is intended to include the unique growth factor isolated and purified from any mammalian tissue or other cells including cell lines. Cell lines include, but are not limited to, glioma-derived cell lines such as C6, hs 683 and GS-9L; glioblastomas such as A-172 and T98G; neuroblastomas such as IMR-32 and SK-N-MC; neurogliomas such as H4; tetromas such as XB-2; astrocytomas such as U-87 MG and U-373 MG; embryonal carcinomas and non-transformed glial or astrocyte cell lines, and the human medulloblastoma line TE 671, with GS-9L and TE 671 being preferred. VEGF AB is present and can be isolated from rat tissue including ovary, heart and kidney. Anterior pituitary tumor cell lines such as GH3 and Hs 199 may also be used. It is intended that VEGF of this invention can be obtained from any mammal species capable of producing VEGF, this includes, but is not limited to, rat and human.

Vascular endothelial cell growth factor may exist in various microheterogeneous forms which are isolated from one or more of the various cells or tissues described above. Microheterogeneous forms as used herein refer to a single gene product, that is a peptide produced from a single gene unit of DNA, which is structurally modified at the mRNA level or following translation. Peptide and protein are used interchangeably herein. The microheterogeneous forms will all have similar mitogenic activities. Biological activity and biologically active are used interchangeably and are herein defined as the ability of VEGF to stimulate DNA synthesis in target cells including vascular endothelial cells as described below which results in cell proliferation. The modifications may take place either in vivo. or during the isolation and purification process. In vivo modification results from, but is not limited to, proteolysis, glycosylation, phosphorylation, deamidation or acetylation at the N-terminus. Proteolysis may include exoproteolysis wherein one or more terminal amino acids are sequentially, enzymatically cleaved to produce microheterogeneous forms which have fewer amino acids than the original gene product. Proteolysis may also include endoproteolytic modification that results from the action of endoproteases which cleave the peptide at specific locations within the amino acid sequence. Similar modifications can occur during the purification process which also results in production of microheterogeneous forms. The most common modification occurring during purification is proteolysis which is generally held to a minimum by the use of protease inhibitors. Under most conditions one or more microheterogeneous forms are present following purification of native VEGFs. Native VEGFs refers to VEGF Isolated and purified from cells that produce VEGFs. Vascular endothelial cell growth factor may also exist in various alternatively spliced forms which is defined herein as the production of related mRNAs by differential processing of exons and introns. Exons are defined as those parts of the DNA sequence of a eukaryotic gene that code for the final protein product. It is also intended that the present Invention includes VEGF subunits A,B and C which are defined as comprising the full length translation products of all alternatively spliced mRNAs made from the gene encoding the subunits and their corresponding mature amino acid sequences generated by proteolytic removal of the amino terminal secretory leader amino acid sequences. It is further intended that the invention only include those microheterogeneous and alternatively spliced VEGF subunits which when in the dimenc form exhibit biological activity such as vascular endothelial cell stimulation as discussed below.

Glioma cells such as the rat cell line GS-9L are grown to confluence in tissue culture flasks, about 175 cm², in a cell culture medium such as Dulbecco's Modified Eagle's Medium (DMEM) supplemented with about 10% newborn calf serum (NCS). When the cells reach confluence the culture medium is removed, the cell layers are washed with Ca⁺⁺, Mg⁺⁺-free phosphate buffered saline (PBS) and are removed from the flasks by treatment with a solution of trypsin, about 0.1%, and EDTA, about 0.04%. The cells, about 1 x 10⁸, are pelleted by centrifugation, resuspended in about 1500 ml of DMEM containing about 5% NCS and plated into a ten level cell factory (NUNC), 6,000 cm² surface area. The cells are incubated for about 48 to about 96 hours, with 72 hours preferred, at about 37° C in an atmosphere of about 5% CO₂. Following incubation the medium is removed and the cell factories are washed about 3 times with PBS. About 1500 ml of fresh culture media is added containing about a 1:2 mixture of Ham's-F12/DMEM containing about 15 mM Hepes, pH about 7.4, about 5 μg/ml insulin, about 10 μg/ml transferrin and with or without about 1.0 mg/ml bovine serum albumin. This medium is replaced with fresh medium after about 24 hr and collected every 48 hr thereafter. The collected conditioned medium is filtered through Whatmen #1 paper to remove cell debris and stored at about -20° C.

The GS-9L conditioned medium is thawed and brought to pH 6.0 with 1 M HCl. The initial purification step consists of cation exchange chromatography using a variety of cation exchangers on a variety of matrices such

as CM Sephadex C-50, Pharmacia Mono S, Zetachrom SP and Polyaspartic Acid WCX (Nest Group) with CM Sephadex C-50 (Pharmacia) being preferred. The VEGF-containing culture medium is mixed with CM Sephadex C-50 at about 2 gm per about 20 L of the conditioned medium and stirred at low speed for about 24 hr at 4° C. The resin is allowed to settle and the excess liquid is removed. The resin slurry is packed into a column and the remaining culture medium is removed. Unbound protein is washed from the column with 0.05 M sodium phosphate, about pH 6.0, containing 0.15 M NaCl. The VEGF AB is eluted with about 0.05 M sodium phosphate, about pH 6.0, containing about 0.6 M NaCl.

The active fractions collected from the CM Sephadex C-50 column are further fractionated by lectin affinity chromatography for additional purification of VEGF AB. The lectins which may bind VEGF AB include, but are not limited to, lectins which specifically bind mannose residues such as concanavalin A and lens culinaris agglutinin, lectins which bind N-acetylglucosamine such as wheat germ agglutinin, lectins that bind galactose or galactosamine and lectins which bind sialic acids, with concanavalin A (Con A) being preferred. A 0.9 cm diameter column containing about 5 ml packed volume of Con A agarose (Vector Laboratories) is washed and equilibrated with about 0.05 M sodium acetate, about pH 6.0, containing about 1 mM CaCl₂, about 1 mM MnCl₂ and about 0.6 M NaCl. The unbound protein is washed from the column with equilibration buffer. The VEGF AB is eluted with about 0.1 M NaCl buffer containing about 0.32 M α-methyl mannoside and about 0.28 M α-methyl glucoside.

The VEGF AB active eluate from the Con-A column is applied to a Polyaspartic Acid WCX cation exchange high performance liquid chromatography (HPLC) column, 4.6 mm x 250 mm, pre-equilibrated in about 0.05 M sodium phosphate buffer, pH 6.0. The column is eluted with a linear gradient of about 0 to 0.75 M NaCl in the phosphate buffer over about 60 minutes. The flow rate is maintained at about 0.75 ml/min collecting 0.75 ml fractions. Vascular endothelial cell growth factor AB activity is present in fractions eluting between approximately 21.7 and 28.5 ml.

The active fractions eluted from the polyaspartic WCX column that contain VEGF AB are pooled, adjusted to about pH 7.0 and loaded onto a 1 x 10 cm column of Pharmacia Chelating Sepharose 6B charged with an excess of copper chloride and equilibrated in about 0.05 M sodium phosphate, about pH 7.0, containing about 2 M NaCl and about 0.5 mM imidazole (A buffer). VEGF AB is eluted from the column with a gradient from 0-20% B over 10 minutes, 20-35% B over 45 minutes and 35-100% B over 5 minutes at a flow rate of 0.3 ml/min, where B buffer is 0.05 M sodium phosphate, pH 7.0, containing about 2 M NaCl and 100 mM imidazole. The active fractions containing VEGF AB activity eluted between about 12.6 and 22.8 ml of the gradient effluent volume.

The pooled fractions containing VEGF AB activity eluted from the metal chelate column are loaded onto a 4.6 mm x 5 cm Vydac

 C_4 reverse phase HPLC column (5 μ m particle size) previously equilibrated in solvent A [0.1% trifluoroacetic acid (TFA)]. The column is eluted with a linear gradient of about 0 to 30% solvent B over 15 minutes, 30% B for an additional 15 minutes, then 30-45% B over 22.5 minutes and finally 45-100% B over 5.5 minutes. Solvent B consists of solvent A containing 67% acetonitrile (v/v). The bow rate is maintained at about 0.75 ml/min and fractions are collected every minute. The homogeneous VEGF AB elutes from the C_4 column under these conditions at between about 32 and about 38 ml of the gradient effluent volume.

Purity of the protein is determined by sodium dodecylsulfate (SDS) polyacrylamide gel electrophoresis (PAGE) in 12.5% crosslinked gels using the technique of Laemmli, Nature 227:680-684 (1970). The silver stained gels show VEGF AB to consist of one band under non-reducing conditions with an approximate apparent molecular mass of about 58,000 daltons. When a sample containing the microheterogeneous forms of VEGF AB is separated under reducing conditions it migrates as two about 23 kilodalton (kDa) subunits. The purification process results in VEGF AB that is essentially free of other mammalian cell products, such as proteins. Recombinantly derived VEGF AB will also be free of mammalian cell products.

Biological activity is determined by mitogenic assay using mammalian vascular endothelial cells. Human umbilical vein endothelial (HUVE) cells are plated on gelatin-coated dishes at a density of about 5000 cells per well in about 500 μ l of Medium 199 (M199) containing about 20% heat-inactivated fetal calf serum (FCS). Samples to be assayed are added at the time of plating. The tissue culture plates are incubated at about 37° C for about 12 hours and about 2 microcuries of tritiated thymidine (NEN, 20 Ci/mmol) is added per ml of assay medium (1.0 μ Ci/well). The plates are incubated for a further 60 hr, the assay medium is removed and the plates are washed with Hanks balanced salt solution containing about 20 mM Hepes, about pH 7.5, and about 0.5 mg/ml bovine serum albumin. The cells are lysed and the labelled DNA solubilized with about 200 μ l of a solution containing about 2 gm of sodium carbonate and about 400 mg sodium hydroxide in about 100 ml water. The incorporated radioactivity was determined by liquid scintillation counting. The concentration of VEGF which elicited a half-maximal mitogenic response in HUVE cells was approximately 2 \pm 1 ng/ml. The glycosaminoglycan heparin, which is required in these assays at a level of 10-100 μ g/ml to promote a response to a positive control, acidic fibroblast growth factor, does not enhance mitogenic stimulation of these cells by VEGF AB.

A purified about 1-2 μg sample of VEGF AB is reduced in about 0.1 M Tris, about pH 9.5, with about 0.1% EDTA, about 6 M guanidinium chloride and about 20 mM dithiothreitol for about 2 hr at about 50°C. The reduced protein is carboxymethylated for about 1 hour in a solution containing about 9.2 μM of unlabelled and 2.8 μM of ¹⁴C-iodoacetic acid in about 0.7 M Tris, about pH 7.8, and about 0.1% EDTA and about 6 M guanidinium chloride. The protein is carboxymethylated for about 1 hr at room temperature. The protein is isolated after reduction and carboxymethylation by reverse phase HPLC chromatography on a Vydac C₄ column, about 4.6 mm x 5 cm. The protein subunits are loaded onto a column pre-equilibrated with about 0.1% TFA and eluted by a 45 ml linear gradient from about 0.1% TFA to 0.1% TFA/67% acetonitrile at a bow rate of about 0.75 ml/min. The reduced and carboxymethylated protein eluted as two peaks at approximately 23 and 25 ml with the proportion being approximately equal as determined by monitoring absorbance at 210 nm.

Samples of the reduced and carboxymethylated monomers are applied to polybrene-coated glass fiber filters and their N-terminal sequences are determined by Edman degradation in an ABI gas phase microsequencer in conjunction with an ABI 120A on line phenylthiohydantoin analyzer following the manufacturers instructions. The protein showing the peak of absorbance eluting at approximately 25 ml (A subunit or monomer) yielded an amino terminal sequence of: SEQ ID NO:1

Ala Pro Thr Thr Glu Glu Glu Glu Lys Ala His Glu Val Val which is identical to the A chain monomers of VEGF AA, Conn et al., Proc. Natl. Acad. Sci. USA <u>87</u>: 2628-2632 (1990). The peak of absorbance eluting at approximately 23 ml (B subunit or monomer) yielded an N-terminal sequence of: SEQ ID NO:2

Ala Leu Ser Ala Gly Asn Xaa Ser Thr Ser Thr Glu Met Glu Val Val Pro Phe Asn Glu Val

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plus a nearly equal amount of a truncated form of the same sequence missing the first three amino acid residues. The missing Xxx residue corresponds to an Asn residue in the cloned cDNA, see below. Since this missing Asn occurs in a classical Asn Xxx Ser/Thr N-glycosylation sequence it is presumed to be glycosylated. The A subunit and the total of both B subunits are recovered in nearly equal amounts supporting the interpretation that the two peptides combine to form an AB heterodimer in VEGF AB.

A sample of the A monomer was treated with either the protease trypsin which cleaves polypeptides on the C-terminal side of lysine and arginine residues or Lys C which cleaves polypeptides on the C-terminal side of lysine by procedures well known in the art. The peptides are isolated by reversed phase - HPLC(RP-HPLC). The amino acid sequences of the isolated peptides are determined using the Edman degradation in the ABI gas phase sequenator in conjunction with the ABI 120 A on line phenylthiohydantoin analyzer following manufacturer's instructions. The amino acid sequences are shown in Figure 1.

Reduced and carboxymethylated A monomer is dried and solubilized in about 0.7 M Tris, about pH 7.8, about 6 M guanidinium chloride containing about 0.1% EDTA. V8 protease is added in 0. 1 M ammonium bicarbonate buffer, about pH 8.0, and the mixture is incubated for about 48 hr at about 37°C. The protease cleaves predominantly on the carboxyl terminal side of glutamic acid residues. The resulting polypeptides were resolved by C₁₈ RP-HPLC as above.

The reduced and carboxymethylated A subunit protein solution is adjusted to a pH of about 6.8 with 6 N HCI and dithiotreitol is added to a final concentration of 2 M for reduction of any methionine sulfoxide to methionine residues. After about 20 hr of reduction at about 39°C the protein is repurified by C₄ HPLC. The product is dried and cleaved on the carboxyl terminal side of methionine residues by 200 µl of 40 mM cyanogen bromide in about 70 % (v/v) formic acid under an argon atmosphere at about 20°C for about 24 hr in the dark. The cleavage products are resolved by C₁₈ RP-HPLC. The amino acid sequence is shown in Figure 1, see Conn et al., Proc. Natl. Acad. Sci USA 87:2628-2632 (1990).

The full length 190 amino acid residue protein translation product of the VEGF AB, A monomer or subunit, which is now known to be identical with the VEGF AA, A monomer, and its cDNA coding sequence are shown in Figures 2 and 6. The mature amino terminus begins at residue 27, immediately following a typical hydrophobic secretory leader sequence. A single potential N-glycosylation site exists at Asn₁₀₀. Most (143 amino acid residues) of the 164 residues of the reduced and carboxymethylated mature subunit including the amino terminus and HPLC reversed phase-purified products of tryptic (T), Lys-C (L), Staphylococcus aureus V8 protease (V8) and cyanogen bromide (CB) cleavages, were determined by direct microsequencing (Applied Biosystems 470A) using a total of 5 µg of protein. All residues identified by amino acid sequencing are denoted by arrows pointing to the right either directly beneath the mature processed sequence following the bracket at residue 27 for the amino terminal determination of the whole subunit or, for residues identified from the polypeptide cleavage products, above the double-headed arrows spanning the length of the particular polypeptide. One listed pair of polypeptides, V18A and V18B, was sequenced as a mixture and, therefore, are only confirmatory of the cDNA-deduced amino acid sequence, see Figures 1 and 5.

Samples of the reduced and carboxymethylated pure VEGF AB, A and B monomers, were each digested

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with the Lys-C endoproteinase, which cleaves polypeptides on the C-terminal side of lysine residues. The peptides were isolated by reverse phase HPLC and their amino acid sequences were determined as described above. The locations of the peptides in the final VEGF AB, A and B sequences are shown in Figure 2 and Figure 3, respectively.

The full length coding region of the A subunit or monomer is determined from three sets of overlapping cDNA clones. Degenerate oligonucleotide primers based on the amino acid sequences Phe-Met-Asp-Val-Tyr-Gin from polypeptide L42 (residues 42-47) and Cys-Lys-Asn-Thr-Asp from polypeptide T38 (residues 164-168) (see Figure 1) were used to PCR amplify the central region of the cDNA for VEGF A chain following the procedure of Saiki et al., Science 230: 1350- 1354 (1985). A single band migrating at 420 bp was gel purified, digested with Sall, ligated into pGEM3Zf(+) and sequenced. The nucleotide sequence obtained (p4238) was used to design antisense and sense PCR primers to amplify the 5' and 3' ends of the cDNA according to the protocol described by Frohman et al. Proc. Natl. Acad. Sci. USA 85:8998-9002 (1988). These 5' and 3' clones are denoted p5-15 and pW3, respectively. Regions of complete DNA sequences, excluding the primers, determined for each set of clones are Indicated by double-headed arrows above the nucleotide sequence. In addition to the cDNA coding the 164 amino acid secreted form identified by protein sequencing, two alternatively spliced cDNAs encoding a 146 amino acid and a 214 amino acid forms are cloned and sequenced, Figures 4, 5 and 6.

The full length coding region of the B subunit or monomer is determined from four sets of overlapping cDNA clones. Degenerate oligonucleotide primers based on the amino acid sequences from polypeptide L50 are used to PCR amplify the central region of the cDNA for VEGF AB, B monomer, following the procedure of Saiki et al., Science 230: 1350-1354 (1985). A single band migrating at 108 bp was gel purified, digested with Sall, ligated into pGEM3Zf(+) and sequenced. The nucleotide sequence obtained (pYG) was used to design antisense and sense PCR primers to amplify the 5' and 3' ends of the cDNA according to the protocol described by Frohman et al. Proc. Natl. Acad. Sci. USA 85:8998-9002 (1988). These 5' and 3' clones are denoted p5V2 and p3V2, respectively. Additional 5' end sequences are determined from clone 202 isolated from a cDNA library prepared from GS-9L poly A+ RNA. Regions of complete DNA sequences, excluding the primers, determined for each set of clones are indicated by double-headed arrows above the nucleotide sequence. The entire base sequence for the 158 amino acid microheterogeneous B subunit and the 138 amino acid microheterogeneous B subunit are shown in Figures 7 and 8.

The full length coding region of the C subunit or monomer is determined from three sets of overlapping cDNA clones. Degenerate oligonucleotide primers based on the amino acid sequence Phe Ser Pro Ser Cys Val and Glu Met Thr Phe Ser Gly from rat VEGF B subunit are used to PCR amplify the central region of the cDNA of VEGF C chain following the procedure of Saiki et al., Science 230: 1350-1354 (1985). A band migrating at 180 bp is gel purified, reamplified and digested with Sall, Ilgated into pGEM3Zf(+) and sequenced. The nucleotide sequence obtained (pFSEM') is used to design antisense and sense PCR primers to amplify the 5' and 3' ends of the cDNA according to the protocol described by Frohman et al., Proc. Natl. Acad. Sci. USA 85: 8998-9002 (1988). The 5' and 3' clones are denoted p5:16 and p3:19, respectively. The entire base sequence and amino acid sequence for the C subunit are shown in Figure 9.

It is intended that vascular endothelial cell growth factor of the present invention exist as a heterodimer consisting of an A microheterogeneous and/or alternatively spliced subunit or a B microheterogeneous and/or alternatively spliced subunit combined with a C microheterogeneous and/or alternatively spliced subunit. It is further intended that VEGF homodimer of the present invention exist as two C subunits. The native forms of the A, B, C subunits may be processed form alternatively spliced full length translation products. The heterodimers or heterodimeric species can be depicted as: A+B, A+C or B+C with the A, B or C subunits existing in any of the alternatively spliced or microheterogeneous forms. The homodimers or homodimeric species can be formed by combinations of any of the alternatively spliced or microheterogeneous forms. It is also intended that the invention include all of the individual subunit forms of the A subunit, the B subunit and the C subunit of VEGF.

It is further intended that the nucleotide sequence for vascular endothelial cell growth factor be interpreted to include all codons that code for the appropriate amino acids in the sequence for each of the vascular endothelial growth factor subunits, as indicated by the degeneracy of the genetic code. It is further intended that the nucleotide sequence and the amino acid sequence for VEGF subunits include truncated genes or proteins that result in proteins which exhibits biological activity similar to vascular endothelial cell growth factor. The scope of the invention is Intended to include all naturally occurring mutations and allelic varients and any randomly generated artifical mutants which may change the sequences but do not alter biological activity as determined by the ability to stimulate the division of vascular endothelial cells.

The above described heterodimers, homodimers and subunits of vascular endothelial cell growth factor are characterized by being the products of chemical synthetic procedures or of procaryotic or eucaryotic host

expression of the DNA sequences as described herein. A monomer is defined as a subunit that is not incorporated in an oligomeric unit. Expression of the recombinant VEGF genes (recombinant DNA) is accomplished by a number of different host cells which contain at least one of a number of expression vectors. Expression vectors are defined herein as DNA sequences that are required for the transcription of cloned copies of recombinant DNA sequences or genes and the translation of their mRNAs in an appropriate host. Such vectors can be used to express genes in a variety of hosts such as bacteria, bluegreen algae, yeast cells, insect cells, plant cells and animal cells, with mammalian cells being preferred. The genes may also be expressed using any of a number of virus expression systems. Specifically designated vectors allow the shuttling of DNA between bacteria-yeast, bacteria-plant or bacteria-animal cells. An appropriately constructed expression vector should contain: an origin of replication for autonomous replication in host cells, selective markers, a limited number of useful restriction enzyme sites, a high copy number, strong promoters and efficient translational stop signals. A promoter is defined as a DNA sequence that directs RNA polymerase to bind to DNA and to initiate RNA synthesis. A strong promoter is one which causes mRNAs to be initiated at high frequency. Expression vectors may include, but are not limited to, cloning vectors, modified cloning vectors, specifically designed plasmids or viruses and cosmids. The expression of mammalian genes in cultured mammalian cells is well known in the art. Sambrook et al., Molecular Cloning, A Laboratory Manual, 2nd Edition, Book 3, Cold Springs Harbor Laboratory Press (1989) and Cwrent Protocols In Molecular Biology, Ausubel et. al. Eds, Greene Publishing Associates and Wiley-Interscience, 1987 and supplements, disclose various mammalian expression vectors and vector systems along with methods for the introduction of recombinant vectors into mammalian cells. The cDNA for the monomemo forms of the A, B and C subunits can be expressed in a system such as that described by Linemeyer et al., European Patent Application, Publication No. 259,953. The cDNA is incorporated into a commercially available plasmid such as pKK 223-3 (Pharmacia) as modified as by Linemeyer et al. and expressed in E. coli. Other expression systems and host cells are well known in the art.

The high Cys content and glycoslyation sites of the A, B and C subunits along with the structure of the homoand heterodimers suggest that expression of biologically active proteins can be carried out in animal cells. Expression may be carried out in Chinese hamster ovary (CHO) cells with the cloned VEGF DNA cotransfected
with the gene encoding dihydrofolate reductase (dhfr) into dhfr- CHO cells, see Sambrook et al. Transformants
expressing dhfr are selected on media lackling nucleosides and are exposed to increasing concentrations of
methotrexate. The dhfr and VEGF genes are thus coamplified leading to a stable cell line capable of expressing
high levels of VEGF. The plasmid is designed to encode either an A subunit, a B subunit or a C subunit or a
combination of any two of these subunits. The two cDNAs are operably attached so that the protein produced
will be dimeric and will have VEGF biological activity. Operably attached refers to an appropriate sequential
arrangement of nucleotide segments, cDNA segments or genes such that the desired protein will be produced
by cells containing an expression vector containing the operably attached genes, cDNA segments or nucleotides. Plasmids containing a single subunit species may be used to cotransfect a suitable cell line.

The expressed proteins (homodimers or heterodimers) are isolated and pwified by standard protein purification processes. It is to be understood that the expression vectors capable of expressing heterodimeric forms of VEGF will contain two DNA sequences which will encode either an A subunit and/or a DNA sequence which will encode a B subunit and/or a DNA sequence which will encode a C subunit. Expression vectors capable of expressing homodimeric forms of VEGF will contain either one or two DNA sequences which encode either two A, two B or two C subunits.

The ability of the various species of VEGF to stimulate the division of vascular endothelial cells makes this protein in all microheterogeneous forms and alternative splicing forms useful as a pharmaceutical agent. The protein as used herein is intended to Include all microheterogeneous forms as previously described. The protein can be used to treat wounds of mammals including humans by the administration of the novel protein to patients in need of such treatment.

The novel method for the stimulation of vascular endothelial cells comprises treating a sample of the desired vascular endothelial cells in a nutrient medium with mammalian VEGF, preferably human or rat, at a concentration of about 1-10 ng/ml. If the vascular endothelial cell growth is conducted in vitro, the process requires the presence of a nutrient mediuin such as DMEM or a modification thereof and a low concentration of calf or bovine serum such as about 0 to 2% by volume. Preservatives such as antibiotics may also be included; these are well known in the art.

The novel growth factors of this invention are useful for the coverage of artificial blood vessels with vascular endothelial cells. Vascular endothelial cells from the patient would be obtained by removal of a small sequent of peripheral blood vessel or capillary-containing tissue and the desired cells would be grown in culture in the presence of VEGF and any other supplemental components that might be required for growth. After growth of adequate numbers of endothelial cells in culture to cover a synthetic polymeric blood vessel the cells would be plated on the inside surface of the vessel, such as fixed umbilical vein, which is then implanted in the patient.

Alternatively, tubular supports are coated in vitro with VEGF prior to implantation into a patient. Following implantation endothelial cells migrate into and grow on the artificial surface. Prior coating of the artificial vessel either covalently or noncovalently, with proteins such as fibrin, collagen, fibronectin or laminin would be performed to enhance attachment of the cells to the artificial surface. The cell-lined artificial vessel would then be surgically implanted into the patient and, being lined with the patients own cells, would be immunologically compatible. The non-thrombogenic endothelial cell lining should decrease the incidence of clot formation on the surface of the artificial vessel and thereby decrease the tendency of vessel blockage or embolism elsewhere.

The novel proteins are also used for the production of artificial vessels. Vascular endothelial cells and smooth muscle cells from the patient would be obtained and grown separately in culture. The endothelial cells would be grown in the presence of VEGF as outlined above. The smooth muscle would be grown in culture by procedures well known in the art. A tubular mesh matrix of a biocompatible polymer (either a synthetic polymer, with or without a coating of proteins, or a non-immunogenic biopolymeric material such as surgical suture thread) would be used to support the culture growth of the smooth muscle cells on the exterior side and vascular endothelial cells on the interior surface. Once the endothelial cells form a confluent monolayer on the inside surface and multiple layers of smooth muscle cells cover the outside, the vessel is implanted into the patient.

The novel peptides can also be used for the induction of tissue repair or growth. The pure VEGF would be used to induce and promote growth of tissue by inducing vascular growth and /or repair. The peptide can be used either topically for tissue repair or intravascularly for vascular repair. For applications involving neovascularization and healing of surface wounds the formulation would be applied directly at a rate of about 10 ng to about 1 mg/cm²/day. For vascular repair VEGF is given intraveneously at a rate of about 1 ng to about 100 µg/kg/day of body weight. For internal vascular growth, the formulation would be released directly into the region to be neovascularized either from implanted slow release polymeric material or from slow release pumps or repeated injections. The release rate in either case is about 10 ng to about 100 µg/day/cm³.

For non-topical application the VEGF is administrated in combination with pharamaceutically acceptable carri ers or diluents such as, phosphate buffer, saline, phosphate buffered saline, Ringer's solution, and the like, in a pharamaceutical composition, according to standard pharamaceutical practice. For topical application, various pharmaceutical formulations are useful for the administration of the active compound of this invention. Such formulations include, but are not limited to, the following: ointments such as hydrophilic petrolatum or polyethylene glycol ointment; pastes which may contain poms such as xanthan gum; solutions such as alcoholic or aqueous solutions; gels such as aluminum hydroxide or sodium alginate gels; albumins such as human or animal albumins; collagens such as human or animal collagens; celluloses such as alkyl celluloses, hydroxyalkyl celluloses, for example methylcellulose, hydroxyethyl cellulose, carboxymethyl cellulose, hydroxypropyl methylcellulose, and hydroxypropyl cellulose; polyoxamers such as Pluronic® Polyols exemplified by Pluronic® F-127; tetronics such as tetronic 1508; and alginates such as sodium alginate.

The following examples illustrate the present invention without, however, limiting the same thereto.

EXAMPLE 1

Preparation of Medium Conditioned By GS-9L Cells

GS-9L cells were grown to confluence in 175 cm² tissue culture flasks in Dulbecco's Modified Eagle's Medium/10% newborn calf serum (DMEM/NCS). At confluence the medium was decanted from the flasks, the flasks were washed with calcium and magnesium free phosphate buffered saline (PBS) and the cells were removed by treatment with a 1X solution of trypsin/EDTA (Gibco). The cells (1 x 10^{4}) were pelleted by centrifugation, resuspended in 1500 ml of DMEM/5% NCS and plated into a ten level (6000 cm² surface area) cell factory (NUNC). After 72 hours incubation at 37° C in a 5% CO₂ atmosphere the medium was decanted and the cell factories were washed 3 times with PBS. The cells were refed with 1500 ml of a 1:2 mixture of Ham's F-12/DMEM containing 25 mM Hepes, pH 7.4, 5 μ g/ml insulin, 10 μ g/ml transferrin and 1.0 mg/ml bovine serum albumln. This medium was changed with fresh F-12/DMEM after 24 hours and collected every 48 hours after that. The conditioned medium was filtered through a Whatman #1 paper to remove cell debris and stored frozen at -20°C.

EXAMPLE 2

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Carboxymethyl-Sephadex Chromatography of VEGF AA and VEGF AB

of CM Sephadex C-50 cation exchange (Pharmacia) resin preequilibrated in PBS adjusted to pH 6.0 with 1 N HCI were added to 20 liters of conditioned medium. The mixture was stirred at low speed for 24 hours at 4° C. The resin was then allowed to settle and the medium was siphoned off. The remaining resin slurry was packed into a 3.0 cm diameter column and any remaining medium was allowed to drain off. Unbound protein was washed off the column with 0.05 M sodium phosphate, pH 6.0, containing 0.15 M NaCl. Vascular endothelial growth factor activity was eluted from the column with a subsequent wash of 0.05 M sodium phosphate, pH 6.0, containing 0.6 M NaCl.

EXAMPLE 3

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Concanavalin A (Con A) Lectin Affinity Chromatography of VEGF AA and VEGF AB

A 0.9 cm diameter column containing about 5 ml of packed Con A agarose (Vector Laboratories) was equilibrated with 0.05 M sodium acetate, pH 6.0, containing 1 mM Ca⁺⁺, 1 mM Mn⁺⁺ and 0.6 M NaCl. The active eluate from the CM Sephadex C-50 column, Example 2, was applied to the Con A agarose and unbound protein was washed from the column with equilibration buffer. The column was then rinsed with three column volumes of 0.05 M sodium acetate, pH 6.0, containing 1 mM Ca⁺⁺, 1 mM Mn⁺⁺ and 0.1 M NaCl. Bound protein was subsequently eluted from the column by application of this buffer supplemented with 0.32 M α-methyl mannoside and 0.28 M α-methyl glucoside.

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EXAMPLE 4

Polyaspartic Acid WCX HPLC Cation Exchange Chromatography of VEGF AA and VEGF AB

The active eluate from the Con A column, Example 3, was applied to a 25 cm x 4.6 mm poly(aspartic acid) WCX cation exchange HPLC column (Nest Group) pre-equilibrated in 0.05 M sodium phosphate buffer, pH 6.0. The column was eluted with a linear gradient of 0 to 0.75 M NaCl in this buffer over 60 minutes at a flow rate of 0.75 ml/min collecting 0.75 ml fractions. VEGF AB activity present in fractions eluting between approximately 21.7 and 28.5 ml were pooled.

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EXAMPLE 5

Metal Chelate Chromatography

The active fractions eluted from the poly(aspartic acid) WCX column, Example 4, that contain VEGF AB were pooled, adjusted to pH 7.0 and loaded onto a 1 x 10 cm column of Pharmacia Chelating Sepharose 6B charged with an excess of copper chloride and equilibrated in 0.05 M sodium phosphate, pH 7.0, containing 2 M NaCl and 0.5 mM imidazole (A buffer). VEGF AB was eluted from the column with a gradient from 0-20% B over 10 minutes, 20-35% B over 45 minutes and 35-100% B over 5 minutes at a flow rate of 0.3 ml/min, where B buffer was 0.05 M sodium phosphate, pH 7.0, containing 2 M NaCl and 100 mM imidazole. The active fractions containing VEGF AB activity eluting between 12.6 and 22.8 ml of the gradient effluent volume were pooled.

EXAMPLE 6

45 Reverse Phase Chromatography

The fractions containing VEGF AB activity pooled from the metal chelate column, Example 5 were loaded onto a 4.6 mm x 5 cm Vydac C_4 reverse phase HPLC column (5 μ m particle size) equilibrated in solvent A (0.1% trifluoroacetic acid (TFA)). The column was eluted with a gradient of 0-30% solvent B over 15 minutes, 30% B for an additional 15 minutes, then 30-45% B over 22.5 minutes and finally 45-100% B over 5.5 minutes where solvent B = A containing 67% acetonitrile. The flow rate was maintained at 0.75 ml/min. The active VEGF AB fractions eluting between approximately 32.2 and 37.5 ml of the gradient effluent volume were pooled.

EXAMPLE 7

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Mitogenic Assays

Human umbilical vein endothelial cells (HUVE) were plated on gelatin-coated 48 well tissue culture dishes

at a density of 5000 cells/well In 500 μ l of Medium 199 containing 20% heat inactivated fetal calf serum (FCS). Samples to be assayed were added at the time of plating. The tissue culture plates are incubated at 37° C for 12 hours and 2 microcuries of tritiated thymidine (NEN, 20 Ci/mmol) was added per ml of assay medium (1.0 μ Ci/vell). The plates were incubated for a further 60 hr, the assay medium was removed and the plates were washed with Hanks balanced salt solution containing 20 mM Hepes, pH 7.5, and 0.5 mg/ml bovine serum albumin. The cells were lysed and the labelled DNA solubilized with 200 μ l of a solution containing 2 gm of sodium carbonate and 400 mg sodium hydroxide in 100 ml water. The incorporated radioactivity was determined by liquid scintillation counting.

The concentration of VEGF AB which elicited a half-maximal mitogenic response in HUVE cells was approximately 2 ± 1 ng/ml. The glycosaminoglycan heparin, which is required in these assays at a level of 10-100 μ g/ml to promote a response to a positive control, acidic fibroblast growth factor, does not enhance mitogenic stimulation of these cells by VEGF AB.

EXAMPLE 8

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Purity And Protein Structural Characterization of VEGF AB

Purity of the protein under non-reducing conditions was determined by SDS-PAGE in 12.5% crosslinked gels according to the method of Laemmli, Nature 227: 680-685 (1970). The silver-stained gel contained a single band with an apparent mass of approximately 58 kDa. VEGF AB migrated in SDS-PAGE under reducing conditions in 15% crosslinked gels as a broad silver-stained band with apparent molecular mass of approximately 23 kDa.

VEGF AB was stored a 4°C in the aqueous trifluoroacetic acid (TFA)/acetonitrile mixture used to elute the homogeneous protein in reversed phase C₄ HPLC chromatography at the final stage of the purification protocol previously described. Aliquots of the purified protein (1-2 μg) were vacuum evaporated to dryness in acid-washed 10 x 75 mm glass tubes and reduced for 2 hours at 50°C in 100 μl of 0. 1 M Tris buffer, pH 9.5, and 6 M guanidinium chloride containing 0.1% EDTA and 20 mM dithiothreitol (Calbiochem, Ultrol grade) under an argon atmosphere. The reduced protein was subsequently carboxymethylated for 1 hour at 20°C by the addition of 100 μl of 0.7 M Tris, pH 7.8, containing 0.1 % EDTA, 6 M guanidinium chloride, 9.2 μM unlabeled iodoacetic acid and 50 μCi of iodo[2-14 C]acetic acid (17.9 mCi/mmole, Amersham). After completion of the carboxymethylation, the mixture was loaded directly onto a 4.6 mm x 5.0 cm Vydac C₄ column which had been preequilibrated in 0.1% TFA. The reduced and carboxymethylated protein was repurified by elution with a 45 minute linear gradient of 0 to 67% (v/v) acetonitrile in 0.1% TFA at a flow rate of 0.75 ml/min and stored in this elution solution at 4°C. The reduced and carboxymethylated protein eluted as two peaks at approximately 23 and 25 ml that were of approximately equal area as determined by monitoring absorbance at 210 nm.

Samples of the two protein subunits isolated after reduction and carboxymethylation were each applied to polybrene-coated glass fiber filters and their N-terminal sequences were determined by Edman degradation in an ABI gas phase microsequencer in conjunction with an ABI 120A on line phenylthiohydantoin analyzer following manufacturers instructions. The peak of absorbance eluting at approximately 25 ml (A subunit) yielded an amino terminal sequence Ala Pro Thr Thr Glu Gly Glu Gln Lys Ala His Glu Val Val SEQ ID NO: 1 identical to VEGF AA. The peak of absorbance eluting at approximately 23 ml (B subunit) yielded the N-terminal sequence Ala Leu Ser Ala Gly Asn Xaa Ser Thr Glu Met Glu Val Val Pro Phe Asn Glu Val SEQ ID NO: 2 plus a nearly equal amount of a truncated form of the same sequence missing the first three residues. The missing X residue corresponds to an Asn in the cloned sequence. Since this missing Asn occurs in a classical Asn-X-Ser/Thr N-glycosylation sequence it is presumed to be glycosylated. The A and sum of the B chain peptides were recovered in nearly equal amounts supporting the interpretation that the two peptides combine to form an AB heterodimer in VEGF II.

Reduced and carboxymethylated A and B subunits (650 ng each) were each dried by vacuum evaporation in acid-washed 10 x 75 mm glass tubes. Lys C protease (50 ng, Boehringer Mannheim), an enzyme that cleaves on the carboxyl terminal side of lysine residues, was added to each tube in 100 μl of 25 mM Tris, pH 8.5, 0.1 % EDTA. The substrate protein subunits were separately digested at 37°C for 8 hours and the resulting polypeptides resolved by reversed phase HPLC chromatography on a 4.6 mm x 25 cm Vydac C₁₈ column equilibrated in 0.1% TFA. Polypeptides were fractionated by elution with a 2 hour linear gradient of 0-67% acetonitrile in 0.1% TFA at a flow rate of 0.75 ml/min at 20°C. Individual peaks were manually collected and stored in this elution solution at 4°C.

The amino acid sequences of the isolated peptides were then determined using Edman degradation in an ABI gas phase sequenator in conjunction with the ABI 120 A on line phenylthichydantoin analyzer (Applied Biosystems Int.). The peptide sequences are shown in the following Figures 2 and 3. The amino acid sequence

of Lys C fragment L20 (Fig. 5) demonstrates that the form of VEGF AB mature A subunit in the heterodimer is the 164 amino acid form. The amino acid sequence of Lys C fragment L26 (Fig. 3) demonstrates that the form of VEGF AB mature B subunit in the heterodimer is the 135 amino acid form derived from the 158 full length amino acid form.

EXAMPLE 9

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Cloning and Sequencing of the VEGF A Monomer

10 PCR Amplification, Cloning and Sequencing of P4238

Two degenerate oligonu leotides were synthesized in order to amplify the cDNA encoding the peptide sequences of VEGF A subunit between LysC fragment L 42 and tryptic fragment T38. These oligonucleotides were:

L42.2 5 TTTGTCGACTT[TC]ATGGA[TC]GT[N]TA[TC]CA 3 SEQ ID NO:3

T383B

5' CAGAGAATTCGTCGACA[AG]TC[N]GT[AG]TT[TC]TT [AG]CA 3' SEQ ID NO:4

where N=ACGT

Poly A* RNA was isolated from GS-9L cells using the Fast Track RNA isolation kit from Invitrogen and the protocol provided. First strand cDNA synthesis was performed as follows;

1 μg of GS-9L RNA was annealled to 1 μg of adapter primer TA17,

5' GACTCGAGTCGACTTTTTTTTTTTTTTTTT 3' SEQ ID NO:5, by incubating in a volume of 10 μl at 70°C for 5 min. followed by cooling to room temperature. To this reaction was added:

- 3.0 µl water
- 2.5 µl 10X buffer (500 mM Tris-HCl pH 8.3, 750 mM KCl, 100 mM MgCl₂, 5 mM spermidine)
- 2.5 µl 100 mM DTT
- 2.5 µl 10 mM each dATP, dGTP, dCTP, dTTP
- s 0.6 بلا 15 units RNasin
 - 2.5 µl 40 mM Na pyrophosphate
 - 1.5 μl 15 units reverse transcriptase

and the reaction was incubated at 42°C for 1 hour, then diluted to 1 ml in 10 mM Tris-HCl I mM EDTA, pH 7.5.

PCR Reactions:

Primary reaction (100 µl)

- 10 μl 10X buffer from Perkin Elmer Cetus GeneAmp kit
- из 16 µl 1.25 mM each stock of dATP, dCTP, dGTP, and dTTP
 - 2 μl first strand GS9L cDNA
 - 2 µJ 50 pMoles L42.2
 - 2 µl 50 pMoles Т383' В
 - 0.5 μl 2.5 units Amplitaq DNA polymerase
- o 67.5 µl water

Reaction conditions, 40 cycles of 94°C, 1'; 50°C, 2'30"; 72°C, 2'.

Prep scale secondary reaction:

- is 100 µl 10X buffer
 - 160 µl 1.25 mM each stock of dATP, dCTP, dGTP, and dTTP
 - 10 µl primary PCR reaction
 - 20 µl 500 pMoles L42.2

20 பி 500 pMoles T383'B

5 μl 25 units Amplitaq DNA polymerase

685 µl water

Reaction conditions 94°C, 1'; 55°C, 2'; 72°C, 2'; 30 cycles.

The PCR product was concentrated by Centricon 30 spin columns, purified on a 1% agarose gel, and digested with restriction endonuclease Sall. The Sall fragment was then ligated into Sall cut pGEM3Zf(+). The ligation mix was used to transform <u>E. coli</u> XL-1 blue. Plasmid DNA was isolated from white transformants and sequenced by the dideoxy chain termination method.

10 PCR Amplification, Cloning and Sequencing of pW-3

Based on the sequence obtained from the p4238 clones, two specific PCR primers were synthesized; oligo 307 5' TTTGTCGACTCAGAGCGGAGAAAGC 3' SEQ ID NO:6 and oligo 289 5' TTTGTCGACGAAAAAT-CACTGTGAGC 3' SEQ ID NO:7. These primers were used in combination with oligoA 17

5'GACTCGAGTCGACATCG 3' SEQ ID NO:8 to amplify the cDNA encoding the COOH terminus of VEGF A subunit using the 3' RACE technique described by Frohman et al., PNAS 85: 8998-9002 (1988).

PCR reactions:

20 Primary reaction 100 ய

10 µi 10X buffer from Perkin Elmer Cetus GeneAmp kit

18 μl 1.25 mM each stock of dATP, dCTP, dGTP, and dTTP

0.35 μl first strand GS-9L cDNA

2 μl 50 pMoles oligo 289

25 0.5 μl 2.5 units Amplitaq DNA polymerase

67.15µl water

Reaction conditions 94°C, 1'; 58°C, 2'; 72°C, 2'; 10 cycles then add 50 pMoles A17, then 1 cycle of 94°C, 1'; 58°C, 2'; 72°C, 40' followed by 40 cycles 94°C, 1'; 58°C, 2'; 72°C, 2'.

30 Prep Scale secondary reaction:

60 μl 10X buffer

108 µl 1.25 mM each stock of dATP, dCTP, dGTP, and dTTP

24 μl primary PCR reaction

i 12 µl 300 pMoles oligo 307

12 µl 300 pMoles oligo A17

3 μl 15 units Amplitaq DNA polymerase

381µl water

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Reaction conditions 94°C, 1'; 58°C, 2'; 72°C, 2'; 30 cycles.

The PCR product was pwified on a 1% agarose gel and digested with restriction endonuclease Sa1I. The Sa1I fragment was then ligated into Sa1I cut pGEM3Zf(+). The ligation mix was used to transform <u>E. coli</u> XL-1 blue. Plasmid DNA was isolated from white transformants and sequenced by the dideoxy chain termination method.

45 PCR Amplification, Cloning and Sequencing of p5-15

Based on the sequence of p4238 clones, two specific PCR primers were synthesized; oligo 113 5'TTTGTCGACACACAGGACGGCTTGAAG 3' SEQ ID NO:9 and oligo 74 5'
TTTGTCGACATACTCCTGGAAGATGTCC 3' SEQ ID NO"10.

These primers were used in combination with oligo A17 5' GACTCGAGTCGACATCG 3' SEQ ID NO:8 to amplify the cDNA encoding the amino terminus of VEGF A subunit using the 5' RACE technique described by Frohman et al., supra. Oligo 151 was synthesized in order to specifically prime VEGF A subunit cDNA from GS-9L RNA. Oligo 151 is 5'

CTTCATCATTGCAGCAGC 3' SEQ ID NO:11.

RNA was isolated from GS-9L cells using the Fast Track RNA isolation kit from Invitrogen using the protocol provided. First strand cDNA synthesis was performed as follows;

One μg of GS9L RNA was annealled to 1 μg of oligo 151 by incubating in a volume of 6 μl at 70°C for 5' followed by cooling to room temperature. To this reaction was added:

- 1.5 μ 10X buffer (500mM Tris-HCl, pH 8.3, 750 mM KCl, 100 mM MgCl₂, 5 mM spermidine)
 2.5 μ 10 mM DTT
 2.5 μ 10 mM each dATP, dGTP, dCTP, dTTP
 0.6 μ 25 units RNasin
 2.5 μ 40 mM Na pyrophosphate
- 9.5 µl 20 units diluted reverse transcriptase

The reaction was incubated at 42°C for 1 hour. Excess oligo151 was removed by Centricon 100 spin columns and the 5' end of the cDNA was tailed by the addition of dATP and terminal transferase. The tailed cDNA was diluted to a final volume of 150 µl in 10 mM Tris-HCl, 1 mM EDTA, pH 7.5.

PCR Reactions:

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Primary reaction (50 μl)

5 山 10X buffer from Perkin Elmer Cetus GeneAmp Kit 8 山 1.25 mM each stock of dATP,dCTP,dGTP, and dTTP 5 山 first strand GS-9L cDNA prime with oligo 151 and tailed 1 山 25 pMoles oligo 113 1 山 25 pMoles oligo A17 1 山 10 pMoles oligo TA17 0.25 山 1.25 units Amplitq DNA polymersase 28.75 山 water

Reaction conditions; 1 cycle 94°C 1'; 50°C 2'; 72°C 40' then 40 cycles of 94°C 1'; 50°C 1'30"; 72°C 2'

Prep scale secondary reaction:

25 60 μl 10X buffer

96 μl 1.25 mM each stock of dATP, dCTP, dGTP, and dTTP

6 µl primary PCR reaction 12 µl 300 pMoles oligo 74 12 µl 300 pMoles oligo A17

3 μl 15 units Amplitaq DNA polymerase

411 µl water

Reaction conditions 94°C, 1'; 55°C, 2'; 72°C, 2' 30 cycles.

The PCR product was concentrated by Centricon 100 spin columns, and digested with restriction endonuclease Sall. The Sall fragment was then ligated into Sall cut pGEM3Zf(+). The ligation mix was used to transform E. coli XL-1 blue. Plasmid DNA was isolated from white transformants and sequenced by the dideoxy chain termination method. The base sequence is shown in Fig. 5.

Cloning and sequencing of alternative forms of VEGF A cDNA

Based on the sequence obtained from the p5-15 and pW-3 clones, two specific PCR primers were synthesized; oligo 5' C 5' TTTGTCGACAACCATGAACTTTCTGC 3' SEQ ID NO:12 and oligo 181 5' TTTGTCGACGGTGAGAGGTCTAGTTC 3' SEQ ID NO:13. These primers were used together to amplify multiple cDNAs encoding alternative forms of the VEGF A subunit.

Preparative PCR Reaction:

50 μl 10X buffer
80 μl 1.25mM each stock of dATP, dCTP, dGTP, and dTTP
10 μl first strand GS-9L cDNA
10 μl 300pMoles oligo 5'C
10 μl 300pMoles oligo 181
2.5 μl 15 units Amplitaq DNA polymerase
337.5 μl water
Reaction conditions 94°C, 1'; 58°C, 2'; 72°C, 3'; 40 cycles.

The PCR product was extracted with phenol/chloroform, concentrated by Centricon 30 spin columns, precipitated by ethanol, and digested with restriction endonuclease Sal I, and ligated into Sall cut pGEM3Zf(+). The ligation mix was used to transform <u>E.coli</u> XL-1 blue. Plasmid DNA was isolated from white transformants

and sequenced by the dideoxy chain termination method. Three sets of clones were identified. Clone#12 encoded the 190 amino acid form of VEGF A subunit identical to that shown in Fig. 1. The 164 amino acid secreted form of VEGF A subunit is that amino acid sequence running continuously from Ala²⁷ to Arg¹⁹⁰. Clone#14 has a 135 base pair deletion between the second base of the Asn¹⁴⁰ codon and the third base of the Arg¹⁸⁴ codon.

This clone thus encodes a 146 aa form of the VEGF A subunit with the conversion of Asn¹⁴⁰ to Lys¹⁴⁰. The 120 amino acid secreted form of VEGF A subunit runs from Ala²⁷ to Asn¹⁴⁰, which becomes Lys¹⁴⁰ and does not begin until Cys¹⁸⁵, this form also finishes at Arg¹⁹⁰, Figure 4. Clone #16 has a 72 base pair insertion between the second and third base of the Asn¹⁴⁰ codon. This clone thus encodes the 214 amino acid form of the VEGF A subunit with the conversion of Asn¹⁴⁰ to Lys¹⁴⁰, Figure 6.

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EXAMPLE 10

Cloning and Sequencing of the VEGF B Subunit

PCR Amplification, Cloning and Sequencing of pYG

Two degenerate oligonucleotides were synthesized in order to amplify the cDNA encoding the peptide sequences of VEGF B on Lys C fragment L50. These oligonucleotides were:

YI 5' TTTGTCGACATA[TC]AT[TCA]GC[N]GA[TC]GA[AG]C 3' SEQ ID NO:14

GC 5' TTTGTCGACTC[AG]TC[AG]TT[AG]CA[AG]CA[N]CC 3' SEQ ID NO:15 where N=ACGT

RNA was Isolated from GS-9L cells using the Fast Track RNA isolation kit from Invitrogen and the protocol provided. First strand cDNA synthesis was performed as follows;

1 μg of GS-9L poly A+RNA was annealied to 1 μg of adapter primer TA17,

- 3.0 µl water
- 2.5 μ l 10X buffer (500 mM Tris-Hd, pH 8.3, 750 mM KCl, 100 mM MgCl₂, 5mM spermidine)
- 2.5 µl 100 mM DTT
- 2.5 µl 10 mM each dATP, dGTP, dCTP, dTTP
- o 0.6 ய 15 units RNasin
 - 2.5 μl 40 mM Na pyrophosphate
 - 1.5 µl 15 units reverse transcriptase

and the reaction was incubated at 42°C for 1 hour, then diluted to 1 ml in 10 mM Tris-HCl, I mM EDTA, pH 7.5.

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PCR Reactions:

Primary reaction (50µl)

| | 5 µl | 10X buffer from Perkin Elmer Cetus GeneAmp kit |
|----|---------|--|
| 40 | 8 μΙ | 1.25 mM each stock of dATP, dCTP, dGTP, and dTTP |
| | 1 μΙ | first strand GS-9L cDNA |
| | 1 μΙ | 50 pMoles oligo YI |
| | 1μ | 50 pMoles oligo GC |
| | 0.25 µl | 1.25 units Amplitaq DNA polymerase |
| | | |

5 33.75 μl water

Reaction conditions, 40 cycles of 94°C, 1'; 50°C, 2'; 72°C, 2'.

Prep scale reaction:

| 60 யி | 10X buffer |
|---------|---|
| 96 µl | 1.25mM each stock of dATP, dCTP, dGTP, and dTTP |
| 12 µ | first strand 659L cDNA |
| 12 µl | 500pMoles oligo YI |
| 12 ш | 500pMoles oligo GC |
| 3 μΙ | 15 units Amplitaq DNA polymerase |
| لىر 405 | water |
| | 96 ப 12 ப 12 ப 12 ப 12 ப 3 பி |

Reaction conditions 94°C, 1'; 50°C, 2'; 72°C, 2' 40 cycles.

The PCR product was concentrated by Centricon 30 spin columns and digested with restriction endonuc-

lease Sall. The Sall fragment was then ligated into Sall cut pGEM3Zf(+). The ligation mix was used to transform <u>E. coli</u> XL-1 blue. Plasmid DNA was isolated from white transformants and sequenced by the dideoxy chain termination method.

5 PCR Amplification, Cloning and Sequencing of p3V2

Based on the sequence obtained from the pYG clones, a specific PCR primer was synthesized; oligo HP 5' TTTGTCGACACCCTAATGAAGTGTC 3' SEQ ID NO:16.

This primer was used in combination with oligo A17 5'

GACTCGAGTCGACATCG 3' SEQ ID NO:8 to amplify the cDNA encoding the COOH terminus of the VEGF B subunit using the 3' RACE technique described by Frohman et al., PNAS 85: 8998-9002 (1988).

Preparative PCR reaction:

- 5 60 μ 10X buffer from Perkin Elmer Cetus Gene Amp Kit 12 μ first strand 659L cDNA 96 μ 1.25 mM each of dATP, dCTP, dGTP, dTTP 12 μ 300 pMoles oligo A17 12 μ 300 pMoles oligo HP
- 3 µl 15units Amplitaq DNA polymerase

405 μl water

Reaction conditions 1 cycle of 94°C, 1′; 58°C, 2′; 72°C, 2′; followed by 40 cycles 94°C, 1′, 58°C, 2′ and 72°C, 2′.

The PCR product was concentrated by Centricon 30 spin columns, precipitated with ethanol and digested with restriction endonuclease Sa1I. The Sa1I fragment was then ligated into Sa1I cut pGEM3Zf(+). The ligation mix was used to transform <u>E</u>. <u>coli</u>. XL-1 blue. Plasmid DNA was isolated from white transformants and sequenced by the dideoxy chain termination method.

PCR Amplification, Cloning and Sequencing of p5V2

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Based on the sequence of pYG clones, two specific PcR primers were synthesized; oligoVL' 5' TTTGTCGACAACAGCGACTCAGAAGG 3' SEQ ID NO: 17 and oligoVS' 5' TTTGTCGACACTGAATATATGAGACAC 3' SEQ ID NO:18. These primers were used in combination with oligo A17

5' GACTCGAGTCGACATCG 3' SEQ ID NO:8 to amplify the cDNA encoding the amino terminus of the VEGF B subunit using the 5' RACE technique described by Frohman et al., supra. Oligo 151 was synthesized in order to prime cDNA from GS-9L RNA Oligo 151 is 5' CTTCATCATTGCAGCAGC 3' SEQ ID NO:11.

Poly A+RNA was isolated from GS9L cells using the Fast Track RNA isolation kit from Invitrogen using the protocol provided. First strand cDNA synthesis was performed as follows:

One μg of GS9L RNA was annealled to 1 μg of oligo 151 by

incubating In a volume of 6 µl at 70°C for 5' followed by cooling

to room temperature. To this reaction was added:

- 1.5 µl 10X buffer (500 mM Tris-HCl, pH 8.3, 750 mM KCl, 100 mM MgCl₂, 5mM spermidine)
- 2.5 µl 10 mM DTT
- 2.5 μl 10 mM each dATP, dGTP, dCTP, dTTP
- 45 0.6 µl 25 units RNasin
 - 2.5 µl 40 mM Na pyrophosphate
 - $9.5~\mu l$ 20 units diluted reverse transcriptase

The reaction was incubated at 42°c for 1 hour.

Excess oligo 151 was removed by Centricon 100 spin columns and the 5' end of the cDNA was tailed by the addition of dATP and terminal transferase. The tailed cDNA was diluted to a final volume of 150 μl in 10 mM Tris-HCl, 1 mM EDTA,pH 7.5

PCR Reactions:

Primary reaction (50 µl)

- 5 μl 10X buffer from Perkin Elmer Cetus GeneAmp Kit
- 8 μl 1.25 mM each stock of dATP,dCTP,dGTP, and dTTP
- 5 μl first strand GS9L cDNA primed with oligo151 and tailed

| 1 µl | 25 pMoles oligo VL' |
|--------|------------------------------------|
| لبر 1 | 25 pMoles oligo A17 |
| 1 ա | 10 pMoles oligo TA17 |
| 0.25 ய | 1.25 units Amplitq DNA polymersase |
| 28 75 | water |

28.75 д

Reaction conditions; 1 cycle 94°C,1'; 58°C, 2'; 72°C, 40' then 40 cycles of 94°C, 1'; 58°C, 2'; 72°C, 2'.

Prep scale secondary reaction

```
100 山
         10X buffer
160 山
         1.25 mM each stock of dATP, dCTP, dGTP, and dTTP
10 µІ
         primary PCR reaction
20 Д
         500 pMoles oligo VS'
20 Д
         300 pMoles oligo A17
5 μΙ
         25 units Amplitaq DNA polymerase
685 µ
         water
```

Reaction conditions 94°C, 1'; 58°C, 2'; 72°C, 2' 30 cycles.

The PCR product was extracted with phenol/chloroform, concentrated by Centricon 30 spin columns, precipitated by ethanol, and digested with restriction endonuclease Sall. The Sall fragment was purified on 4% Nu-Sieve Agarose gel then ligated into Sall cut pGEM3Zf(+). The ligation mix was used to transform E. coli XL-1 blue. Plasmid DNA was isolated from white transformants and sequenced by the dideoxy chain termination method.

PCR Amplification, Cloning and Sequencing of pCV2 and pCV2.1

25

Based on the sequences of the p3V2 and p5CV2 clones, two specific PCR primers were synthesized; oligo 5'CV2.1

5' TTTGTCGAC[N][N]GCAGGTCCTAGCTG 3' SEQ ID NO;19 and oligo 3'CV2 5'

TTTGTCGAC[N][N]CTAATAAATAGAGGG 3' SEQ ID NO:20.

These primers were used together to amplify the cDNA encoding the VEGF 8 subunit.

Preparative PCR Reaction:

```
40 ш
         10X buffer
64 µl
         1.25 mM each dATP, dTTP, dGTP, dCTP
8 μΙ
         first strand GS-9L cDNA
8 ш
         200 pMoles 5'CV2.1
8 μΙ
         200 pMoles 3'CV2
2 μ
         10units Amplitaq DNA polymerase
270 山
```

Reaction conditions: 94°C, 1', 58°C, 2', 72°C, 2'; 40 cycles.

The PCR product was extracted with phenol/chloroform, concentrated by Centricon 30 spin columns, precipitated by ethanol, and digested with restriction endonuclease Sal 1, and ligated into Sal I cut pGEM3Zf(+). The ligation mix was used to transform E. coli XL-1 blue. Plasmid DNA was isolated form white transformants and sequenced by the dideoxy chain termination method. Two sets of clones were identified, one encoded a 158 amino acid sequence and the other encoded a 138 amino acid sequence, see Figures 7 and 8.

cDNA Cloning of VEGF B Subunit

The DNA and protein sequences for the amino terminus of the signal peptide of VEGF B was determined from a cDNA clone isolated from a cDNA library constructed from GS-9L polyA+ RNA.

First Strand Synthesis

50

Anneal 15.6µl (5ug) GS-9L polyA+ RNA and 2.5µl (2.5ug) oligo dT-Xbal primer by heating to 70° C 5' slow cool to room temperature. Add the following:

```
5.5µl
          10X buffer (500 mM Tris-HCl, pH 8.3 (42° C), 750 mM KCl, 100 mM MgCl<sub>2</sub>, 5mM spermidine
5.5யி
          10 mM each dATP, dTTP, dCTP, dGTP
5.5µl
1.4µ1
          (55units) RNasin
```

5.5_ய 40mM NaPPi

13.5_µl (55units) reverse transcriptase

incubate at 42° C 60'.

5 Second Strand Synthesis:

Assemble reaction mix

50 µl first strand reaction

25 µJ 10X buffer (500 mM Tris-HCl, pH7.2, 850 mM KCL, 30 mM MgCl₂ 1mg/ml BSA, 100 mM (NH₄)₂S0₄

7.5 பி 100 mM DTT

25 µl 1mM NAD

6.5 µl (65units) E. coli DNA Polymerasel

2.5 µl (2.5units) E. coli DNA Ligase

2.5 µl (2 units) E. coli RNase H

ร 135 ม water

20

30

50

Incubate at 14° C for 2h and then incubate 70° C for 10′. Add 1ul (10 units) T4 DNA Polymerase, incubate at 37° C for 10′, add 25 μ 0.2M EDTA an extract with phenol/chloroform, then precipitate by the addition of 0.5 volume of 7.5 M ammonium acetate and 3 volumes of ethanol, collect precipitate and resuspend in 20 μ l of 10 mM Tris-HCl, pH 7.5, 1mM EDTA.

cDNA Library Construction

The above cDNA was ligated into EcoR1/ Xbal digested LambdaGEM-4 (Promega Biochemicals) after the addition of EcoR1 linkers and digestion with EcoR1 and Xbal. A cDNA library was amplified from ~50, 000 independent clones.

Isolation of Rat VEGF B cDNA Clone

25 The above cDNA library was screened by placque hybridization using pCV2 as a probe. Hybridization conditions were as follows:

5XSSC (1XSSC is 0.15M sodium chloride, 0.015M sodium citrate,

50% Formamide

5X Denhardt's Solution (1% Ficoll, 1% polyvinylpyrrolidone, 1% bovine serum albumin)

0.15 mg/ml salmon sperm DNA hybridize overnight at 42° C.

Filters were washed 3 times in 2XSSC, 0.1% SDS at room temerature for 5', then 1 time in 1XSSC, 0.1% SDS at 50C for 30'. Positive clones were identified by autoradiography.

The DNA from phage #202 was digested with restriction endonuclease Spel and the 1.1kb band ligated into Xbal digested pGEM3Zf(+). The ligation mix was used to transform E.coli XL-1 blue. Plasmid DNA was isolated from white transformants and sequenced by the dideoxy chain termination method. The cDNA sequence and predicted amino acid sequence of the signal peptide are shown in Figures 7 and 8.

The entire nucleotide and amino acid sequence of the 138 amino acid form is shown in Fig. 7. The secreted protein starts at Ala²⁴ and continues to Arg¹³⁸. The entire nucleotide and amino acid sequence of the 158 amino acid form is shown in Figures 8. The secreted protein starts at Ala²⁴ and continues to Leu¹⁵⁸.

EXAMPLE 11

Cloning and sequencing VEGF C Subunit

5 PCR Amplification, Cloning and Sequencing of pFSEM

Two degenerate oligonucleotides were synthesized based on the sequence of rat VEGF B monomer in order to amplify VEGF cDNAs from the human medulloblastoma line TE-671, ATCC HTB (McAllister et al., Int. J. Cancer 20:206-212 [1977]). These oligonucleotides were:

FS 5'TTTGTCGACA TTC AGT CC(N) TC(N) TG(TC) GT 3' SEQ ID NO:21

EM' 5' TITGTCGACA CTG AGA GAA (N)GT CAT (CT)TC 3'

SEQ ID NO:22

where N= AGCT

Poly A+ RNA was isolated from TE-671 cells using the Fast Track RNA isolation kit from Invitrogen and the protocol provided. First strand cDNA synthesis was performed as follows using the cDNA Cycle kit from Invitrogen;

1μl 1μg of TE-671 polyA+ RNA

19µl water

5μ 100mM MeMgOH 6.25μ 0.7M B-mercaptoethanol 2.5μ random primer 2.5μ RNase Inhibitor 10μ 5X RT buffer 2.5μ 25mM dNTPs 1.25μ reverse transcriptase 12.5units

The reaction was incubated for 60' at 42°C, then 3' at 95°C, placed on ice, then an additional 1.25ul reverse transcriptase was added and the reaction incubated an additional 60' at 42°C.

The above procedure was performed in duplicate and the cDNAs pooled to a final volume of 100ul.

PCR Reactions:

10

Primary reaction (100µl)

15 10 ய 10X buffer from Perkin Elmer Cetus GeneAmp kit 16 ய 1.25mM each of dATP, dCTP, dGTP, TTP

10 μl first strand TE-671 cDNA 2 μl 50 pmoles FS primer 2 μl 50 pmoles EM' primer

2 μl 50 pmoles EM' primer 0 0.5 μl 2.5 units Amplitaq DNA polymerase

59.5 µl water

Reaction conditions: 40 cycles of 90°C, 1'; 2' ramp to 45°C; 2' at 45°C; 2' at 72°C.

Gel Purification

 $20~\mu$ l of the primary PCR reaction was purified on a 4% NuSieve agarose gel. The 180 base pair band was excised from the gel. heated to 65°C for 5' and used directly as template for the secondary PCR reaction.

Secondary PCR reaction 200µl

20 μl 10X buffer from Perkin Elmer Cetus GeneAmp kit

32 ய 1.25mM each of dATP. dCTP, dGTP, TTP

5 μl melted gel slice

4 μl 100 pmoles FS primer

4 μl 100 pmoles EM' primer

1 μl 5 units Amplitaq DNA polymerase

134 µl water

Reaction conditions: 35 cycles of 94°C, 1'; 50°C, 2'; 72°C, 2'

The PCR product was purified on a Qiagen tip 5 column, then digested with restriction endonuclease Sall. The Sall fragment was then ligated into Sall cut pGEM3Zf(+), and the ligation mix used to transform <u>E. coli</u> XL-1 blue. Plasmid DNA was isolated from white transformants and sequenced by the dideoxy chain termination method.

40 PCR Amplification, Cloning and Sequencing of p3'.19

Based on the sequence obtained from the pFSEM' clone, a specific PCR primer was synthesized; oligo LH 5' TTTGTCGACA CTG CAC TGT GTG CCG GTG 3' SEQ ID NO:23. This primer was used in combination with oligo A17,5' GACTCGAGTCGACATCG 3' SEQ ID NO:24, to amplify the cDNA encoding the COOH terminus of the VEGF C subunit using the 3' RACE technique described by Frohman et al., PNAS 85:8998-9002 (1988).

Poly A+ RNA was isolated from TE-671 cells using the Fast Track RNA isolation kit from Invitrogen and the protocol provided. First strand cDNA synthesis was performed as follows using the cDNA Cycle kit from Invitrogen and the TA17 adapter primer.

50 TA17 5' GACTCGAGTCGACATCGATTTTTTTTTTTTTT 3' SEQ ID NO:5

0.8 μl 1μg of TE-671 polyA+ RNA

20.7 µl water

5 山 100 mM MeMgOH

6.25 µl 0.7 M B-mercaptoethanol

5 1.0 ய 0.88 µg primer TA17

2.5 山 RNase Inhibitor

10 μl 5X RT buffer

2.5 لبر 25mM dNTPs

1.25 山 reverse transcriptase 12.5 units

The reaction was incubated for 60' at 42°C, then 3' at 95°C, placed on ice, then an additional 1.25ul reverse transcriptase was added and the reaction incubated an additional 60' at 42°C.

3' RACE PCR

- 20 ш 10 X buffer from Perkin Elmer Cetus GeneAmp kit 1.25mM each of dATP, dCTP, dGTP, TTP 32 µl 20 ш first strand TE-671 cDNA primed with TA17 2 μ 50 pmoles LH primer 50 pmoles A17 primer 2 Д 5 units Amplitag DNA polymerase 1.0 山

123 д

15

Reaction conditions: 40 cycles of 94C, 1'; 2' at 58°C; 3' at 72°C.

The PCR product was purified on a Qiagen tip 5 column, then digested with restriction endonuclease Sall. The Sall fragment was then ligated into Sall cut pGEM3Zf(+), and the ligation mix used to transform E. coli XL-1 blue. Plasmid DNA was isolated from white transformants and sequenced by the dideoxy chain termination method.

PCR Amplification, Cloning and Sequencing of p5'.16

Based on the sequence obtained from the pFSEM' clone, two specific PCR primers were synthesized; oligo VE' 5' TTTGTCGACA AC ATT GGC CGT CTC CAC C 3' SEQ ID NO:24, and oligo TG' 5' TTTGTCGACA ATC GCC GCA GCC GGT 3' SEQ ID NO:25. These primers were used in combination with

oligo A17, 5' GACTCGAGTCGACATCG 3' SEQ ID NO:8, and oligo TA17

5'GACTCGAGTCGACATCGATTTTTTTTTTTTTTTT3' SEQ ID NO:5 to amplify the cDNA encoding the amino terminus of the VEGF C subunit using the 5' RACE technique described by Frohman et al., PNAS 85: 8998-9002 (1988).

Poly A+ RNA was isolated from TE-671 cells using the Fast Track RNA isolation kit from Invitrogen and the protocol provided. First strand cDNA synthesis was performed as follows using the cDNA Cycle kit from Invitrogen and the VE' primer.

```
1 μg of TE-671 polyA+ RNA
1.0 µl
الر 20.25
             water
5 μΙ
             100 mM MeMgOH
6.25 µl
             0.7 M B-mercaptoethanol
             1.0 µg primer VE'
1.0 山
2.5 д
             RNase Inhibitor
10 µ
             5X RT buffer
2.5 山
             25 mM dNTPs
0.5 ய
             AMV reverse transcriptase (Promega) 10units
```

The reaction was incubated for 60'at 42°C, then 3' at 95°C, placed on ice, then an additional 1.25ul reverse transcriptase was added and the reaction incubated an additional 60' at 42°C. Excess oligo VE' was removed by a Centricon 100 spin column and the 5' end of the cDNA was tailed by the addition of dATP and terminal transferase. The tailed cDNA was diluted to a final volume of 200 ul in 10mM Tris-HCl, 1mM EDTA, pH 7.5.

5' RACE PCR 5 X 100ul

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10X buffer from Perkin Elmer Cetus GeneAmp kit 10 山 16 ш 1.25mM each of dATP, dCTP, dGTP, TTP 10 д first strand TE-671 cDNA primed with VE' 2 ш 50 pmoles TG' primer 2 ш 50 pmoles A17 primer 2 ш 20 pmoles TA17 primer 0.5 ப 2.5 units AMplitaq DNA polymerase 57.5 µl water

Reaction conditions: 40 cycles of 94°C, 1'; 2' ramp to 58°C; 2' at 58°C; 2' at 72°C.

The PCR product was purified on a Qiagen tip 5 column, then digested with restriction endonuclease Sall. The Sall fragment was then ligated into Sall cut pGEM3Zf(+), and the ligation mix used to transform E. coli

XL-1 blue. Plasmid DNA was isolated from white transformants and sequenced by the dideoxy chain termination method. The combined sequences form plasmids pFSEM', p3'19 and p5'16 are shown in Figure 9.

PCR Amplificaotion, Cloning and Sequencing of phVC16 and phVC2

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Based on the sequences of the p5'. 16 and p3'. 19 clones, two specific PCR primers were synthesized; oligo 5' GCVB 5' TTTGTCGAC TGG CTC TGG ACG TCT GAG 3' SEQ ID NO:26 and oligo 3'VC 5'TTTGTCGAC ACT GAA GAG TGT GAC GG 3' SEQ ID NO:27. These primers were used together to amplify the cDNA encoding the complete VEGF C subunit.

Poly A+ RNA was isolated from TE-671 cells using the Fast Track RNA isolation kit from Invitrogen and the protocol provided. First strand cDNA synthesis was performed as follows using the cDNA Cycle kit from Invitrogen;

```
0.8μl 1μg of TE-671 polyA+ RNA
19.2μl water
5μl 100 mM MeMgOH
```

6.25µl 0.7 M B-mercaptoethanol

2.5山 oligo dT primer 2.5山 RNase Inhibitor 10山 5X RT buffer 2.5山 25 mM dNTPs

1.25µl reverse transcriptase 12.5units

The reaction was incubated for 60' at 42C, then 3' at 95C, placed on ice, then an additional 1.25ul reverse transcriptase was added and the reaction incubated an additional 60' at 42C.

PCR Reaction 200 ul

25 20μl 10X buffer from Perkin Elmer Cetus GeneAmp kit 32μl 1.25mM each of dATP. dCTP, dGTP, TTP 20μl first strand TE-671 cDNA primed with oligo dT 4μl 50 pmoles 5' GCVB primer 4μl 50 pmoles 3'VC primer
 30 1μl 5 units Amplitaq DNA polymerase water

Reaction conditions: 40 cycles of 94°C, 1′;; 2′ at 50°C; 2′at 72°C.

The PCR product was purified on a Qiagen tip 5 column, then digested with restriction endonuclease Sall. The Sall fragment was then ligated into Sall cut pGEM3Zf(+), and the ligation mix used to transform <u>E. coli</u> XL-1 blue. Plasmid DNA was isolated from white transformants and sequenced by the dideoxy chain termination method. In the sequences of clones phVC16 and phVC2 base 463 (Fig. 9) was changed from a T to a C eliminating the translational stop codon following amino acid 154; this results in the addition of 16 amino acids following amino acid Lys 154. The nucleotide sequence and the deduced amino acid sequence of this addition is:

CAG AGA CCC ACA GAC TGC CAC CTG TGC GGC GAT GCT GTT Gln Arg Pro Thr Asp Cys His Leu Cys Gly Asp Ala Val

CCC CGG AGG TAA Pro Arg Arg 170

.70 SEQID NO:29

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In addition clone phVC16 contains a 3 base pair deletion (Figure 9, nucleotide residues 73-75) resulting in the deletion of Gln 25.

| | SEQUENCE | LISTING | | |
|----|----------|-------------------------------|---|--|
| 10 | . (2) | (i) SEQU (A) (B) (C) | ON FOR SEQ ID NO:1: ENCE CHARACTERISTICS: LENGTH: 14 amino acid: TYPE: amino acid STRANDEDNESS: Not App: TOPOLOGY: linear | |
| 15 | | | | |
| | | | | |
| 20 | | | | |
| 25 | | | | |
| 30 | | | | |
| 35 | | | | |
| 40 | | | | |

| | | | (x: | i) | SEQU | UENCE DESCRIPTION: SEQ ID NO:1 | |
|----|-------|-------|-------|----------|------------|---|---|
| | Ala | Pro 3 | Thr I | Chr | | Gly Glu Gln Lys Ala His Glu Val | |
| 5 | | | | | 5 | 10 | |
| | Val | | | | | | |
| | | | | | | | • |
| | (2) | TNF | יבאאר | רז∩א | FOR | R SEQ ID NO:2: | |
| | (2) | 1111 | (i | | | ENCE CHARACTERISTICS: | |
| 10 | | | , | | _ | LENGTH: 19 amino acids | |
| | | | | 1 | (B) | TYPE: amino acid | |
| | | | | | (C) | STRANDEDNESS: Not Applicable | |
| | | | | | | TOPOLOGY: linear | |
| | | | | | | QUENCE DESCRIPTION: SEQ ID NO:2: | |
| 15 | Ala | Leu : | Ser 1 | Ala | GŢĀ | Asn Xaa Ser Thr Glu Met Glu Val | |
| | **- 1 | D 1 | n | . | 21 | 10 | |
| | val | Pro 1 | Pne . | ASN | GIU | val | |
| | | 15 | | | | | |
| 20 | (2) | INF | ORMA' | TION | FOE | R SEQ ID NO:3: | |
| | ,-, | | (i | | | JENCE CHARACTERISTICS: | |
| | | | | | (A) | LENGTH: 26 base pairs | |
| | | | | | (B) | TYPE: nucleic acid | |
| | | | | | | STRANDEDNESS: single | |
| 25 | | | | | (D) | | - |
| | | | | | | QUENCE DESCRIPTION: SEQ ID NO:3: CT TYATGGAYGT NTAYCA 26 | |
| | | | 1.1 | 101 | JOAC | of Italignatus Minion 20 | |
| | (2) | INF | ORMA | TION | 1 FO | OR SEQ ID NO:4: | |
| 30 | | | (i | .) : | SEQU | JENCE CHARACTERISTICS: | |
| | | | | | (A) | | |
| | | | | | | TYPE: nucleic acid | |
| | | | | | (C) (D) | STRANDEDNESS: single TOPOLOGY: linear | |
| | | | / > | | | QUENCE DESCRIPTION: SEQ ID NO:4: | |
| 35 | | | | | | TC GTCGACARTC NGTRTTYTTR CA 32 | |
| | | | | | | | |
| | (2) | INF | ORMA | | | OR SEQ ID NO:5: | |
| | | | (j | L) | SEQU | UENCE CHARACTERISTICS: | |
| 40 | | | | | | LENGTH: 35 base pairs | |
| | | | | | | TYPE: nucleic acid STRANDEDNESS: single | |
| | | | | | | STRANDEDNESS: single TOPOLOGY: linear | |
| | | | () | ei) | SEC | QUENCE DESCRIPTION: SEQ ID NO:5: | |
| | | | GZ | CTC | GAGT | C GACATCGATT TTTTTTTTT TTTTT 35 | |
| 45 | | | | | | | |
| | (2) | INE | ORMA | | | OR SEQ ID NO:6: | |
| | | | (: | i) | - | UENCE CHARACTERISTICS: | |
| | | | | | (A) | | |
| 50 | | | | | (B) | | |
| | | | | | (C) | STRANDEDNESS: single | |
| | | | | | | | |

| 5 | | (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6: TTTGTCGACA ACACAGGACG GCTTGAAG 28 |
|----|-----|---|
| | (2) | INFORMATION FOR SEQ ID NO:7: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: nucleic acid |
| 10 | | (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: TTTGTCGACG AAAATCACTG TGAGC 25 |
| 15 | (2) | INFORMATION FOR SEQ ID NO:8: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single |
| 20 | | (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8: GACTCGAGTC GACATCG 17 |
| 25 | (2) | INFORMATION FOR SEQ ID NO:9: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear |
| 30 | | (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9: TTTGTCGACA ACACAGGACG GCTTGAAG 28 |
| 35 | (2) | INFORMATION FOR SEQ ID NO:10: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10: |
| 40 | | TTTGTCGACA TACTCCTGGA AGATGTCC 28 |
| 45 | (2) | INFORMATION FOR SEQ ID NO:11: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: sizele |
| | | (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11: CTTCATCATT GCAGCAGC 18 |
| 50 | | |

| 5 | (2) | INFORMATION FOR SEQ ID NO:12: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12: TTTGTCGACA ACCATGAACT TTCTGC 26 |
|----|-----|--|
| 15 | (2) | INFORMATION FOR SEQ ID NO:13: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single |
| | | (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13: TTTGTCGACG GTGAGAGGTC TAGTTC 26 |
| 20 | (2) | INFORMATION FOR SEQ ID NO:14: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single |
| 25 | (2) | (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14: TTTGTCGACA TAYATHGCNG AYGARC 26 |
| 30 | (2) | INFORMATION FOR SEQ ID NO:15: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear |
| 35 | | (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15: TTTGTCGACT CRTCRTTRCA RCANCC 26 |
| 40 | (2) | (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear |
| 45 | | (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16: TTTGTCGACA CACCCTAATG AAGTGTC 27 |
| | (2) | INFORMATION FOR SEQ ID NO:17: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid |
| 50 | | (B) TYPE: nucleic acid(C) STRANDEDNESS: single |

| | | (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17: TTTGTCGACA ACAGCGACTC AGAAGG 26 |
|------------|-----|--|
| 5 | | INFORMATION FOR SEQ ID NO:18: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single |
| 10 | | (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18: TTTGTCGACA CTGAATATAT GAGACAC 27 |
| 15 | (2) | INFORMATION FOR SEQ ID NO:19: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear |
| 20 | | (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19: TTTGTCGACN NGCAGGTCCT AGCTG 25 |
| 25 | (2) | INFORMATION FOR SEQ ID NO:20: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear |
| 30 | | (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20: TTTGTCGACN NCTAATAAAT AGAGGG 26 |
| 35 | (2) | (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear |
| 40 | | (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21: TTTGTCGACA TTCAGTCCNT CNTGYGT 27 |
| | (2) | INFORMATION FOR SEQ ID NO:22: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid |
| 4 5 | | (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22: TTTGTCGACA CTGAGAGAAN GTCATYTC 28 |
| 50 | | |

| 5 | (2) | INFORMATION FOR SEQ ID NO:23: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23: TTTGTCGACA CTGCACTGTG TGCCGGTG 28 |
|----|-----|--|
| 10 | (2) | INFORMATION FOR SEQ ID NO:24: |
| 15 | (2) | (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24: TTTGTCGACA ACATTGGCCG TCTCCACC 28 |
| | | 11101COACA ACATIGOCCO ICICCACC 20 |
| 20 | (2) | INFORMATION FOR SEQ ID NO:25: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single |
| 25 | | (D) TOPOLOGY: linear |
| _ | | (xi) SEQUENCE DESCRIPTION: SEQ ID NO:25: TTTGTCGACA ATCGCCGCAG CAGCCGGT 28 |
| | (2) | |
| 30 | | (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear |
| 35 | | (xi) SEQUENCE DESCRIPTION: SEQ ID NO:26: |
| • | | TTTGTCGACT GGCTCTGGAC GTCTGAG |
| | (2) | (i) SEQUENCE CHARACTERISTICS: |
| 40 | | (A) LENGTH: 26 base pairs(B) TYPE: nucleic acid |
| | | <pre>(C) STRANDEDNESS: single (D) TOPOLOGY: linear</pre> |
| | | (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27: |
| 45 | | TTTGTCGACA CTGAAGAGTG TGACGG 26 |
| | (2) | INFORMATION FOR SEQ ID NO:28: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 51 base pairs |
| 50 | | (B) TYPE: nucleic acid |
| ~ | | (C) STRANDEDNESS: single |

| | (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28: |
|----|--|
| 5 | CAG AGA CCC ACA GAC TGC CAC CTG TGC GGC GAT GCT GTT 39 |
| 10 | CCC CGG AGG TAA 51 |
| | (2) INFORMATION FOR SEQ ID NO:29: (1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 16 |
| 15 | (B) TYPE: amino acids (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29: |
| 20 | Gln Arg Pro Thr Asp Cys His Leu Cys Gly Asp Ala Val |
| | 5 10 Pro Arg Arg |
| 25 | 15 (2) INFORMATION FOR SEQ ID NO:30: |
| | (i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 577 base pairs(B) TYPE: nucleic acid |
| 30 | (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30: |
| | AACC 4 |
| 35 | ATG AAC TIT CTG CTC TCT TGG GTG CAC TGG ACC CTG GCT TTA CTG 49 Met Asn Phe Leu Leu Ser Trp Val His Trp Thr Leu Ala Leu Leu 5 10 15 |
| 40 | CTG TAC CTC CAC CAT GCC AAG TGG TCC CAG GCT GCA CCC ACG ACA 94 Leu Tyr Leu His His Ala Lys Trp Ser Gln Ala Ala Pro Thr Thr 20 25 30 |
| | GAA GGG GAG CAG AAA GCC CAT GAA GTG GTG AAG TTC ATG GAC GTC 139 Glu Gly Glu Gln Lys Ala His Glu Val Val Lys Phe Met Asp Val 35 40 40 |
| 45 | TAC CAG CGC AGC TAT TGC CGT CCG ATT GAG ACC CTG GTG GAC ATC 184 Tyr Gln Arg Ser Tyr Cys Arg Pro Ile Glu Thr Leu Val Asp Ile 50 55 60 |
| 50 | TTC CAG GAG TAC CCC GAT GAG ATA GAG TAT ATC TTC AAG CCG TCC 229 Phe Gln Glu Tyr Pro Asp Glu Ile Glu Tyr Ile Phe Lys Pro Ser 65 70 75 |

| 5 | TGT Cys | GTG Val | CCC Pro | CTA Leu | ATG Met 80 | CGG Arg | TGT Cys | GCG Ala | GGC Gly | TGC Cys 85 | TGC Cys | AAT Asn | GAT Asp | GAA Glu | GCC Ala 90 | 274 |
|----|------------|------------|------------|-------------|-------------------|------------|------------|--------------|------------|-------------------|----------------------|------------|------------|------------|-------------------|-----|
| | CTG Leu | GAG Glu | TGC Cys | GTG Val | CCC Pro 95 | ACG Thr | TCG Ser | GAG Glu | AGC Ser | AAC Asn 100 | GTC Val | ACT Thr | ATG Met | CAG Gln | ATC Ile 105 | 319 |
| 10 | Met | Arg | Ile | Lys | Pro 110 | His | Gln | Ser | Gln | His 115 | Ile | Gly | Glu | Met | Ser 120 | |
| 15 | Phe | CTG Leu | Gln | His | Ser 125 | Arg | Суз | Glu | Суз | Arg 130 | Pro | Lys | Lys | qeA | Arg 135 | |
| 20 | ACA Thr | AAG Lys | CCA Pro | GAA Glu | AAT Asn 140 | CAC His | TGT Cys | GAG Glu | CCT Pro | TGT Cys 145 | TCA Ser | GAG Glu | CGG Arg | AGA Arg | AAG Lys 150 | 454 |
| 20 | CAT His | TTG Leu | TTT Phe | GTC Val | CAA Gln 155 | GAT Asp | CCG Pro | CAG Gln | ACG Thr | TGT Cys 160 | AAA Lys | TGT Cys | TCC Ser | TGC Cys | AAA Lys 165 | 499 |
| 25 | AAC Asn | ACA Thr | GAC Asp | TCG Ser | CGT Arg 170 | TGC Cys | AAG Lys | GCG Ala | AGG Arg | CAG Gln 175 | CTT Leu | GAG Glu | TTA Leu | AAC Asn | GAA Glu 180 | 544 |
| 30 | CGT Arg | ACT Thr | TGC Cys | AGA Arg | TGT Cys 185 | GAC Asp | AAG Lys | CCA Pro | AGG Arg | CGG Arg 190 | TGA | | | | | 577 |
| | (2) | II | NFOR | MAT: (i) | SI () | EQUE A) | NCE LEN | CHA GTH: | RAC' | TERI 90 a | ISTI amin cids | o a | cids | : | | |
| 35 | | | | (xi | 1) | C) O) | STR. | ANDE OLOG | DNE Y: | SS: lir | si near | ngle | | МО | : 31 : | |
| 40 | | Asn | | | 5 | | | | | 10 | | | | | 15 | |
| | | Tyr | | | 20 | | | | | 25 | | | | | 30 | - |
| 45 | | Gly | | | 35 | | | · | | 40 | | | | | 45 | |
| | | Gln | | | 50 | | | | | 55 | | | | | 60 | |
| 50 | Phe | G1n | Glu | Tyr | Pro | Азр | Glu | Ile | Glu | Tyr | Ile | Phe | Lys | Pro | Ser | |

| | | | | | 65 | | | | | 70 | | | | | 75 | |
|------------|------------|------------|------------|-------------|------------------|------------|------------|----------------------|-------------|------------------|----------------------|------------|------------|------------|------------------|-----|
| · 5 | Суз | Val : | Pro | Leu : | Met 80 | Arg | Суз | Ala | Gly | Суз 85 | Суз | Asn | Азр | Glu | Ala 90 | |
| | Leu | Glu | Cys | Val | Pro 95 | Thr | Ser | Glu | Ser | Asn 100 | Val | Thr | Met | Gln | 11e 105 | |
| 10 | Met | Arg | Ile | | Pro 110 | His | Gln | Ser | Gln | His 115 | Ile | Gly | Glu | Met | Ser 120 | |
| | Phe | Leu | Gln | | Ser 125 | Arg | Суз | Glu | Cys | Arg 130 | Pro | Lys | Lys | Asp | Arg 135 | |
| 15 | Thr | Lys | Pro | Glu | Asn 140 | His | Суs | Glu | Pro | Cys 145 | Ser | Glu | Arg | Arg | Lys 150 | |
| | His | Leu | Phe | Val | Gln 155 | Asp | Pro | Gln | Thr | Cys 160 | Lys | Суз | Ser | Cys | Lys 165 | |
| 20 | Asn | Thr | Asp | Ser | Arg 170 | Cys | Lys | Ala | Arg | Gln 175 | Leu | Glu | Leu | Asn | Glu 180 | |
| | Arg | Thr | Суз | Arg | Cys 185 | Asp | Lys | Pro | Arg | Arg 190 | | | | | | |
| 25 | (2) | IN | IFOR | MAT: (i) | S) (. | EQUE A) | NCE | CHA GTH | ARAC : 4 | TER | ISTI base c ac | pa | irs | | | |
| 30 | | | | (xi | (| D) | TOP | ANDI OLOG E DI | GY: | li | near | | |) NO | : 32 : | |
| | ACC | A | | | | | | | | | | | | | | 4 |
| 35 | ATG Met | AAC Asn | TTT Phe | CTG Leu | CTC Leu 5 | TCT Ser | TGG Trp | GTG Val | CAC His | TGG Trp 10 | ACC Thr | CTG Leu | GCT Ala | TTA Leu | CTG Leu 15 | 49 |
| 40 | | TAC Tyr | | | | | | | | | | | | | ACA Thr 30 | 94 |
| 45 | GAA Glu | GGG Gly | GAG Glu | CAG Gln | AAA Lys 35 | GCC Ala | CAT | GAA Glu | GTG Val | GTG Val 40 | Lys | TTC Phe | ATG Met | GAC Asp | GTC Val 45 | 139 |
| 45 | TAC Tyr | CAG Gln | CGC | AGC Ser | TAT Tyr 50 | Cys | CGT | CCG | ATT | GAG Glu 55 | Thr | CTG Leu | GTG Val | GAC Asp | ATC Ile 60 | 184 |
| 50 | TTC Phe | CAG Gln | GAG Glu | TAC Tyr | CCC | GAT Asp | GAG | ATA Ile | GAG Glu | TAT Tyr | ATC | TTC Phe | AAG Lys | CCG | TCC Ser | 229 |

| | | | | | 65 | | | | | 70 | | | | | 75 | | |
|----|------------|------------|------------|-------------|-------------------|------------|------------|------------|-------------|----------------------|---------------|------------|------------|------------|-------------------|-------------------|---|
| 5 | TGT Cys | GTG Val | CCC Pro | CTA Leu | ATG Met 80 | CGG Arg | TGT Cys | GCG Ala | G1A GCC | TGC Cys 85 | TGC Cys | AAT Asn | GAT Asp | GAA Glu | GCC Ala 90 | 274 | |
| 10 | CTG Leu | GAG Glu | TGC Cys | GTG Val | CCC Pro 95 | ACG Thr | TCG Ser | GAG Glu | AGC Ser | AAC Asn 100 | GTC Val | ACT Thr | ATG Met | CAG Gln | ATC Ile 105 | 319 | |
| .~ | ATG Met | CGG Arg | ATC Ile | AAA Lys | CCT Pro 110 | CAC His | CAA Gln | AGC Ser | CAG Gln | CAC His 115 | ATA Ile | GGA Gly | GAG Glu | ATG Met | AGC Ser 120 | 364 | |
| 15 | | | | | AGC Ser 125 | | | | | | | | | | | 409 | |
| 20 | | | | | AAA Lys 140 | | | | | | | TGA | | | | 445 | |
| 25 | (2) | I | NFOF | TAMS (i) | () | EQUE | NCE LEN | CHA GTH | ARAC : 1 | :33: TER: 46 a | IST I amir | | .cid | s | | | |
| | | | | (xi | (1 | C) D) | STR | AND: | EDNE GY: | SS: | s: nea: | | | D NC | :33 | : | • |
| 30 | | | | | Leu 5 | | | | | 10 | | | | | 15 | | |
| 35 | | | | | His 20 Lys | | | | | 25 | | | | | 30 | 1 | |
| | | | | | 35 Tyr 50 | Суз | | | | 40 | Thr | | | | 45 |) ! | |
| 40 | Phe | Gln | Glu | Туг | Pro 65 | Asp | Glu | Ile | Glu | | Ile | Phe | E Ly: | s Pro | 5 Se 1 | : 5 | - |
| 45 | | | | | Met 80 | | • | | | 85 | , · | | | | 9(|) | |
| | | | | | Pro 95 Pro | • | | | | 100 |) | | | | 10 | 5 | • |
| 50 | Met | , | , | | 110 | | | | | 115 | | | | | 12 | 0 | |

| | Phe | Leu | Gln | His | Ser 125 | Arg | Суз | Glu | Суз | Arg 130 | Pro | Lys | Lys | Asp | Arg 135 | | |
|-------------|-----|------------|------|-------------|------------------------|-----|---------------------------|-------------------|-------------------------------|--|----------------------------------|-------------------|-----|-------|-------------------|----------|--|
| 5 | Thr | Lys | Pro | Glu | Lys 140 | Cys | Ąsp | Lys | Pro | Arg 145 | Arg | | | | | | |
| 10 | (2) | IN | IFOR | MAT] (i) | SE (<i>I</i> (E | QUE | NCE LEN TYP: STR | ANDE OLOG } | RAC' 6 nuc DNE Y: | TERI 49 t leic SS: lir ENCI | ease ac si near E DE | pai id ngle | 2 ' | : NO: | SEC |) | |
| 15 | | _ | | | | | | • | ID N | 0:3 | 4: | | | | | | |
| | AAC | | | | | | | | | | | | | | | 4 | |
| 20 | _ | AAC Asn | | | - | | | | | | | | | | | 49 | |
| | | TAC Tyr | | | | | | | | | | | | | | 94 | |
| 25 | | GGG Gly | | | | | _ | _ | | | Lys | | | | GTC Val 45 | 139 | |
| <i>30</i> | | CAG Gln | | | | | | | | | | | | | ATC Ile 60 | 184 | |
| 35 | | CAG Gln | | | | | | | | | | | | | TCC Ser 75 | 229 | |
| | | GTG Val | | | | | | | | | | | | | GCC Ala 90 | 274 | |
| 40 | | GAG Glu | | | | | | | | | | | | | ATC Ile 105 | 319 | |
| 45 . | | CGG Arg | | | | | | | | | Ile | | | | AGC Ser 120 | 364 | |
| | | CTG Leu | | | | | | | | | Pro | | | | AGA Arg 135 | 409 | |
| 50 | ACA | AAG | CCA | GAA | AAA | AAA | TCA | GTT | CGA | GGA | AAG | GGA | AAG | GGT | CAA | 454 | |

| | Thr | Lys | Pro | Glu | Lys 140 | Lys | Ser | Val | Arg | Gly 145 | Lys | Gly | Lys | Gly | Gln 150 | |
|----------------|---------------------------------|---------------------------------|--|---|---|--------------------------------|--|--|---|---|--|--------------------------------|---------------------------------|---|--|-----|
| | AAA | CGA | AAG | CGC | DAG | 277 | TCC | ccc | ጥጥጥ | 444 | TCC | TCC | NCC | CTT | CNC | 400 |
| . 5 | Lvs | Arg | Lvs | Ara | Lvs | Lvs | Ser | Ara | Phe | Lve | Sar | Trn | Sar | Val | ui. | 433 |
| | | 5 | -,, 0 | | 155 | -,0 | 561 | mry | * | 160 | 361 | 11p | 3et | Val | 165 | |
| | TGT | GAG | CCT | TGT | TCA | GAG | CGG | AGA | AAG | CAT | TTG | TTT | GTC | CAA | GAT | 544 |
| | Cys | Glu | Pro | Cys | Ser | Glu | Arg | Arg | Lys | His | Leu | Phe | Val | Gln | Asp | |
| 10 | | | | | 170 | | | | | 175 | | | | | 180 | |
| | CCG | CAG | ACG | TGT | AAA | TGT | TCC | TGC | AAA | AAC | ACA | GAC | TCG | CGT | TGC | 589 |
| | Pro | Gln | Thr | Суз | | Суз | Ser | Суз | Lys | Asn | Thr | Asp | Ser | Arg | Суз | |
| | | | | | 185 | | | | | 190 | | | | | 195 | |
| 15 | AAG | GCG | »GG | CNG | ርጥሞ | CXC | ም ሞ እ | 220 | C3.3 | | | mcc | | | | 634 |
| | Lvs | Ala | Ara | Gln | Leu | Glu | LON | AAC | GAA | Ara | The | TGC | AGA | TGT | DAC | 634 |
| | -1- | | | | 200 | 014 | Dea | 11311 | 014 | 205 | 1111 | Cys | vrâ | Cys | 210 | |
| | | | | | | | | | | | | | | | | |
| | | CCA | | | TGA | | | | | | | | | | | 649 |
| 20 | rys | Pro | Arg | Arg | | | | | | | | | | | | |
| | (2) | TI | JEAD | MATT | T / NT | EOD | CE/ | ` TD | | . 25. | | | | | | |
| | (2) | 11 | 1FOR | (i) | | | | | | | | 00. | | | | |
| | | | | (1) | | | | | | | STI | | | _ | | |
| | | | | | - | A) B) | | GTH: | | | | o a | clas | 5 | | |
| 0.5 | | | | | | | | | | | | | | | | |
| 25 | | | | | • | | | | | o a | | 201 | _ | | | |
| 25 | | | | | i | C) | STR | ANDI | EDNE | ss: | si | ngl | е | | | |
| 25 | | | | (xi | (1 | C) D) | STR TOP | ANDI OLO(| EDNE SY: | SS: | si near | : - | | NO. | . 25. | |
| 25 | | | | (xi | (1 | C) D) | STR TOP | ANDI OLO(| EDNE SY: | SS: | si near | : - | |) NO | : 35 : | |
| | Met | Asn | Phe | | () ()) (| C) D) SEQU | STR TOP ENC | ANDI OLOG E DE | EDNE SY: SSCR | SS: lia IPT: | si near ION: | SE | Q IE | | | |
| . 30 | Met | Asn | Phe | | () ()) (| C) D) SEQU | STR TOP ENC | ANDI OLOG E DE | EDNE SY: SSCR | SS: lia IPT: | si near ION: | SE | Q IE | | | |
| | | | | Leu | () () Leu 5 | C) D) SEQU Ser | STR TOP ENC: | ANDI OLO(E DE Val | EDNE SY: SSCR His | SS: lin IPT: Trp 10 | si near ION: Thr | SE(| Q II | Leu | Leu 15 | |
| | | Asn Tyr | | Leu | (())) Leu 5 | C) D) SEQU Ser | STR TOP ENC: | ANDI OLO(E DE Val | EDNE SY: SSCR His | SS: lin IPT: Trp 10 | si near ION: Thr | SE(| Q II | Leu | Leu 15 | |
| | | | | Leu | () () Leu 5 | C) D) SEQU Ser | STR TOP ENC: | ANDI OLO(E DE Val | EDNE SY: SSCR His | SS: lin IPT: Trp 10 | si near ION: Thr | SE(| Q II | Leu | Leu 15 | |
| 30 | Leu | Tyr | Leu | Leu | Leu 5 His 20 | C) D) SEQU Ser Ala | STR TOP ENC: Trp | ANDI OLO(E DE Val | EDNE GY: SSCR His | SS: lin IPT: Trp 10 Gln 25 | sinear ION: Thr | SE(Leu Ala | Q ID | Leu Thr | Leu 15 Thr 30 | |
| | Leu | | Leu | Leu | Leu 5 His 20 | C) D) SEQU Ser Ala | STR TOP ENC: Trp | ANDI OLO(E DE Val | EDNE GY: SSCR His | SS: lin IPT: Trp 10 Gln 25 | sinear ION: Thr | SE(Leu Ala | Q ID | Leu Thr | Leu 15 Thr 30 Val | |
| 30 | Leu | Tyr | Leu | Leu | Leu 5 His 20 | C) D) SEQU Ser Ala | STR TOP ENC: Trp | ANDI OLO(E DE Val | EDNE GY: SSCR His | SS: lin IPT: Trp 10 Gln 25 | sinear ION: Thr | SE(Leu Ala | Q ID | Leu Thr | Leu 15 Thr 30 | |
| 30 | Leu Glu | Tyr | Leu Glu | Leu His Gln | Leu 5 His 20 Lys 35 | C) D) SEQU Ser Ala | STR TOP ENC Trp Lys | ANDI OLOG E DE Val Trp Glu | EDNE GY: SCR His Ser Val | SS: lin IPT: Trp 10 Gln 25 Val 40 | sinear ION: Thr Ala | SE(Leu Ala Phe | O II Ala Pro Met | Leu Thr Asp | Leu 15 Thr 30 Val 45 | |
| 30 | Leu Glu | Tyr Gly | Leu Glu | Leu His Gln | Leu 5 His 20 Lys 35 | C) D) SEQU Ser Ala | STR TOP ENC Trp Lys | ANDI OLOG E DE Val Trp Glu | EDNE GY: SCR His Ser Val | SS: lin IPT: Trp 10 Gln 25 Val 40 | sinear ION: Thr Ala | SE(Leu Ala Phe | O II Ala Pro Met | Leu Thr Asp | Leu 15 Thr 30 Val 45 | |
| 30 | Leu Glu Tyr | Tyr Gly Gln | Leu Glu Arg | Leu His Gln Ser | Leu 5 His 20 Lys 35 | C) D) SEQU Ser Ala Ala Cys | STR TOP ENC: Trp Lys His | ANDI OLOG E DE Val Trp Glu | EDNE GY: ESCR His Ser Val | SS: lin IPT: Trp 10 Gln 25 Val 40 Glu 55 | sinear ION: Thr Ala Lys | SE(Leu Ala Phe Leu | Ala Pro Met | Leu Thr Asp | Leu 15 Thr 30 Val 45 Ile 60 | |
| 30 | Leu Glu Tyr | Tyr Gly | Leu Glu Arg | Leu His Gln Ser | Leu S His 20 Lys 35 Tyr 50 | C) D) SEQU Ser Ala Ala Cys | STR TOP ENC: Trp Lys His | ANDI OLOG E DE Val Trp Glu | EDNE GY: ESCR His Ser Val | SS: lin IPT: Trp 10 Gln 25 Val 40 Glu 55 | sinear ION: Thr Ala Lys | SE(Leu Ala Phe Leu | Ala Pro Met | Leu Thr Asp | Leu 15 Thr 30 Val 45 Ile 60 Ser | |
| 30 | Leu Glu Tyr | Tyr Gly Gln | Leu Glu Arg | Leu His Gln Ser | Leu 5 His 20 Lys 35 | C) D) SEQU Ser Ala Ala Cys | STR TOP ENC: Trp Lys His | ANDI OLOG E DE Val Trp Glu | EDNE GY: ESCR His Ser Val | SS: lin IPT: Trp 10 Gln 25 Val 40 Glu 55 | sinear ION: Thr Ala Lys | SE(Leu Ala Phe Leu | Ala Pro Met | Leu Thr Asp | Leu 15 Thr 30 Val 45 Ile 60 | |
| 30 | Leu Glu Tyr Phe | Tyr Gly Gln Gln | Leu Glu Arg Glu | Leu His Gln Ser | ((((())))))))))))))))))))))))))))))))) | C) D) SEQU Ser Ala Ala Cys | STR TOP ENC: Trp Lys His Arg | ANDI OLOO E DE Val Trp Glu | EDNE GY: CSCR His Ser Val Ile | SS: lin IPT: Trp 10 Gln 25 Val 40 Glu 55 Tyr 70 | sinear ION: Thr Ala Lys | SE(Leu Ala Phe Leu Phe | Pro Met Val | Leu Thr Asp Asp | Leu 15 Thr 30 Val 45 Ile 60 Ser 75 | |
| 30 | Leu Glu Tyr Phe | Tyr Gly Gln | Leu Glu Arg Glu | Leu His Gln Ser | ((((())))))))))))))))))))))))))))))))) | C) D) SEQU Ser Ala Ala Cys | STR TOP ENC: Trp Lys His Arg | ANDI OLOO E DE Val Trp Glu | EDNE GY: CSCR His Ser Val Ile | SS: lin IPT: Trp 10 Gln 25 Val 40 Glu 55 Tyr 70 | sinear ION: Thr Ala Lys | SE(Leu Ala Phe Leu Phe | Pro Met Val | Leu Thr Asp Asp | Leu 15 Thr 30 Val 45 Ile 60 Ser 75 | |
| 30 35 40 | Leu Glu Tyr Phe Cys | Tyr Gly Gln Gln Val | Leu Glu Arg Glu Pro | Leu His Gln Ser Tyr | ((((())))))))))))))))))))))))))))))))) | C) D) SEQU Ser Ala Ala Cys Asp | STR TOP ENC: Trp Lys His Arg Glu | ANDI OLOC E DE Val Trp Glu Pro | EDNE GY: CSCR His Ser Val Ile Glu Gly | SS: lin IPT: Trp 10 Gln 25 Val 40 Glu 55 Tyr 70 Cys 85 | sinear ION: Thr Ala Lys Thr | SE(Leu Ala Phe Leu Phe Asn | Ala Pro Met Val Lys | Leu Thr Asp Asp Pro Glu | Leu 15 Thr 30 Val 45 Ile 60 Ser 75 Ala 90 | |
| 30 | Leu Glu Tyr Phe Cys | Tyr Gly Gln Gln | Leu Glu Arg Glu Pro | Leu His Gln Ser Tyr | ((((())))))))))))))))))))))))))))))))) | C) D) SEQU Ser Ala Ala Cys Asp | STR TOP ENC: Trp Lys His Arg Glu | ANDI OLOC E DE Val Trp Glu Pro | EDNE GY: CSCR His Ser Val Ile Glu Gly | SS: lin IPT: Trp 10 Gln 25 Val 40 Glu 55 Tyr 70 Cys 85 Asn | sinear ION: Thr Ala Lys Thr | SE(Leu Ala Phe Leu Phe Asn | Ala Pro Met Val Lys | Leu Thr Asp Asp Pro Glu | Leu 15 Thr 30 Val 45 Ile 60 Ser 75 Ala 90 | |
| 30 35 40 | Leu Glu Tyr Phe Cys | Tyr Gly Gln Gln Val | Leu Glu Arg Glu Pro | Leu His Gln Ser Tyr | ((((())))))))))))))))))))))))))))))))) | C) D) SEQU Ser Ala Ala Cys Asp | STR TOP ENC: Trp Lys His Arg Glu | ANDI OLOC E DE Val Trp Glu Pro | EDNE GY: CSCR His Ser Val Ile Glu Gly | SS: lin IPT: Trp 10 Gln 25 Val 40 Glu 55 Tyr 70 Cys 85 | sinear ION: Thr Ala Lys Thr | SE(Leu Ala Phe Leu Phe Asn | Ala Pro Met Val Lys | Leu Thr Asp Asp Pro Glu | Leu 15 Thr 30 Val 45 Ile 60 Ser 75 Ala 90 | |
| 30 35 40 | Leu Glu Tyr Phe Cys | Tyr Gly Gln Gln Val | Leu Glu Arg Glu Pro Cys | Leu His Gln Ser Tyr Leu Val | (((((()))))))))))))))))))))))))))))))) | C) D) SEQU Ser Ala Ala Cys Asp | STR TOP ENC: Trp Lys His Arg Glu Cys | ANDI OLOO E DE Val Trp Glu Pro Ile Ala | EDNE GY: CSCR His Ser Val Ile Glu Gly Ser | SS: lin IPT: Trp 10 Gln 25 Val 40 Glu 55 Tyr 70 Cys 85 Asn 100 | sinear ION: Thr Ala Lys Thr Ile Cys | SE(Leu Ala Phe Leu Phe Asn Thr | Pro Met Val Lys Asp | Leu Thr Asp Asp Pro Glu Gln | Leu 15 Thr 30 Val 45 Ile 60 Ser 75 Ala 90 Ile 105 | |
| 30 35 40 | Leu Glu Tyr Phe Cys | Tyr Gly Gln Gln Val | Leu Glu Arg Glu Pro Cys | Leu His Gln Ser Tyr Leu Val | (((((()))))))))))))))))))))))))))))))) | C) D) SEQU Ser Ala Ala Cys Asp | STR TOP ENC: Trp Lys His Arg Glu Cys | ANDI OLOO E DE Val Trp Glu Pro Ile Ala | EDNE GY: CSCR His Ser Val Ile Glu Gly Ser | SS: lin IPT: Trp 10 Gln 25 Val 40 Glu 55 Tyr 70 Cys 85 Asn 100 | sinear ION: Thr Ala Lys Thr Ile Cys | SE(Leu Ala Phe Leu Phe Asn Thr | Pro Met Val Lys Asp | Leu Thr Asp Asp Pro Glu Gln | Leu 15 Thr 30 Val 45 Ile 60 Ser 75 Ala 90 Ile 105 | |

| | Phe | Leu | Gln | His | Ser 125 | Arg | Суз | Glu | Cys | Arg 130 | Pro | Lys | Lys | Asp | Arg 135 | |
|----|-----|-----|------------|------------|------------|-----|-------------------|------------|-------------|--------------|----------------------|-----|-----|------|------------------|-----|
| 5 | Thr | Lys | Pro | Glu | Lys 140 | Lys | Ser | Val | Arg | Gly 145 | Lys | Gly | Lys | Gly | Gln 150 | |
| | Lys | Arg | Lys | Arg | Lys 155 | Lys | Ser | Arg | Phe | Lys 160 | Ser | Trp | Ser | Val | His 165 | |
| 10 | Cys | Glu | Pro | Суз | Ser 170 | Glu | Arg | Arg | Lys | His 175 | Leu | Phe | Val | Gln | Asp 180 | |
| | Pro | Gln | Thr | Суз | Lys 185 | Суз | Ser | Суз | Lys | Asn 190 | Thr | Asp | Ser | Arg | Cys 195 | |
| 15 | Lys | Ala | Arg | Gln | Leu 200 | Glu | Leu | Asn | Glu | Arg 205 | Thr | Cys | Arg | Cys | Asp 210 | |
| - | Lys | Pro | Arg | Arg | | | | | | | | | | | | |
| 20 | (2) | I | NFOR | TAM (i) | \$1 () | | NCE LEN TYP | CHA GTH | ARAC : 4 | TER: 17 l | ISTI base c ac | pa | | | | |
| 25 | | | | (xi | Ċ | D) | TOP | OLO | GY: | 11: | near | : - | |) NO | :36: | |
| 30 | | | GCC Ala | | | | | | | | | | | | | 45 |
| | | | GCT Ala | | | | | | | | | | | | | 90 |
| 35 | | | GAA Glu | | | | | | | | | | | | CGC Arg 45 | 135 |
| 40 | | | TGC Cys | | | | | | | | | | | | GAA Glu 60 | 180 |
| | | | AAT Asn | | | | | | | | | | | | CTT Leu 75 | 225 |
| 45 | | | CGC Arg | | | Gly | | Суз | Gly | Asp | Glu | | | His | TGT Cys | 270 |

| | GTG Val | GCG Ala | CTA Leu | AAG Lys | ACA Thr 95 | GCC Ala | AAC Asn | ATC Ile | ACT Thr | ATG Met 100 | CAG Gln | ATC Ile | TTA Leu | AAG Lys | ATT Ile 105 | 315 |
|----------|------------|------------|------------|-------------|----------------------|------------------------------|---------------------------------|-------------------------------------|-------------------------|----------------------------------|-------------------------------------|------------|------------|------------|-------------------|-----|
| 5 | CCC | CCC Pro | AAT Asn | CGG Arg | GAT Asp 110 | CCA Pro | CAT His | TCC Ser | TAC Tyr | GTG Val 115 | GAG Glu | ATG Met | ACA Thr | TTC Phe | TCT Ser 120 | 360 |
| 10 | CAG Gln | GAT Asp | GTA Val | CTC Leu | TGC Cys 125 | GAA Glu | TGC Cys | AGG Arg | CCT Pro | ATT Ile 130 | CTG Leu | GAG Glu | ACG Thr | ACA Thr | AAG Lys 135 | 405 |
| | | GAA Glu | | TAA | | | | | | | | | | | | 417 |
| 15 20 | (2) | II | NFOR | MAT (i) | SI () () () | EQUE A) B) C) D) | NCE LEN TYP STR TOP | CHA GTH: E: & ANDE OLOC | RAC min DNE Y: | TER: 38 a o a SS: li | ISTI emin cid si sinear | ngl | | | : 37 : | |
| 25 | | Leu | | | 5 | | | | | 10 | | | | | 15 | |
| | Gly | Leu | Ala | Val | His 20 | Ser | Gln | Gly | Ala | Leu 25 | Ser | Ala | Gly | Asn | neA 30 | |
| 30 | Ser | Thr | Glu | Met | Gl u 35 | Val | Val | Pro | Phe | Asn 40 | Glu | Val | Trp | Gly | Arg 45 | |
| | Ser | Tyr | Cys | Arg | Pro 50 | Met | Glu | Lys | Leu | Val 55 | Tyr | Ile | Ala | Asp | Glu 60 | |
| 35 | His | Pro | Asn | Glu | Val 65 | Ser | His | Ile | Phe | Ser 70 | Pro | Ser | Cys | Val | Leu 75 | |
| | Leu | Ser | Arg | Суз | Ser 80 | Gly | Суз | Cys | Gly | Asp 85 | Glu | Gly | Leu | His | Cys 90 | |
| 40 | Val | Ala | Leu | Lys | Thr 95 | Ala | Asn | Ile | Thr | Met 100 | Gln | Ile | Leu | Lys | Ile 105 | |
| | Pro | Pro | Asn | Arg | Asp 110 | Pro | His | Ser | Tyr | Val 115 | Glu | Met | Thr | Phe | Ser 120 | |
| 45 | Gln | Asp | Val | Leu | Cys 125 | Glu | Суз | Arg | Pro | Ile 130 | Leu | Glu | Thr | Thr | Lys 135 | |
| | Ala | Glu | Arg | | | | | | | | | | | | | |
| 50 | (2) | I | NFOR | TAM: (i) | | | _ | _ | | | ISTI | cs: | | | | |

| (A) LENGTH: 477 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:38: | | | | | | | | | | | | | | | | |
|---|-----|---|-----|-------------|-----|------|------|------|------|------|-----|-----|-------|-----|-------------------|-----|
| 5 | | | | (xi |) 5 | SEQU | ENC | E DE | SCR | IPTI | ON: | SE(|) ID | NO | : 38 : | |
| | | | | ATG Met | | | | | | | | | | | | 45 |
| 10 | | | | GTG Val | | | | | | | | | | | | 90 |
| 15 | | | | ATG Met | | | | | | | | | | | CGC Arg 45 | 135 |
| 20 | | | | CGG Arg | | | | | | | | | | | GAA Glu 60 | 180 |
| | | | | GAA Glu | | | | | | | | | | | CTT Leu 75 | 225 |
| 25 | | | | TGT Cys | | | | | | | | | | | TGT Cys 90 | 270 |
| 30 | | | | AAG Lys | | | | | | | | | | | ATT Ile 105 | 315 |
| | | | | CGG Arg | | | | | | | | | | | TCT Ser 120 | 360 |
| 35 | | | | CTC Leu | | | | | | | | | | | AAG Lys 135 | 405 |
| 40 | | | Arg | Arg | Lys | Thr | Lys | Gly | Lys | Arg | Lys | Gln | Ser | Lys | ACC Thr 150 | |
| 45 | | | | GAG Glu | | Pro | | | | | | | | | | 477 |
| _ | (2) | I | NFO | TAMS (i) | S | EQUI | ENCE | CH | ARAC | TER | IST | | .cid: | s | | |

| 5 . | | | | (xi | (C (E | ;))) | STRA TOP(| ANDE OLOG | | SS: lin | si ear | | | NO | : 39 : | |
|-----|-----|------|------|-------------|------------|------------------------|--------------|--------------|----------------------------|------------|--------------|-----------|-----|------|------------------|----|
| | Met | Leu | Ala | Met | Lys 5 | Leu | Phe | Thr | Cys | Phe 10 | Leu | Gln | Val | Leu | Ala 15 | |
| 10 | Gly | Leu- | Ala | Val | His 20 | Ser | Gln | Gly | Ala | Leu 25 | Ser | Ala | Gly | Asn | Asn 30 | |
| | Ser | Thr | Glu | Met | Glu 35 | Val | Val | Pro | Phe | Asn 40 | Glu | Val | Trp | Gly | Arg 45 | |
| 15 | Ser | Tyr | Суз | Arg | Pro 50 | Met | Glu | Lys | Leu | Val 55 | Tyr | Ile | Ala | Asp | Glu 60 | |
| | His | Pro | Asn | Glu | Val 65 | Ser | His | Ile | Phe | Ser 70 | Pro | Ser | Суз | Val | Leu 75 | |
| 20 | Leu | Ser | Arg | Cys | Ser 80 | Gly | Суз | Суз | Gly | Asp 85 | Glu | Gly | Leu | His | Cys 90 | |
| 25 | Val | Ala | Leu | Lys | Thr 95 | Ala | Asn | Ile | Thr | Met 100 | Gln | Ile | Leu | Lys | Ile 105 | |
| | Pro | Pro | Asn | Arg | Asp 110 | Pro | His | Ser | Tyr | Val 115 | Glu | Met | Thr | Phe | Ser 120 | |
| 30 | Gln | Asp | Val | Leu | Cys 125 | Glu | Суз | Arg | Pro | Ile 130 | Leu | Glu | Thr | Thr | Lys 135 | |
| | Ala | Glu | Arg | Arg | Lys 140 | | Lys | Gly | Lys | Arg 145 | Lys | Gln | Ser | Lys | Thr 150 | |
| 35 | Pro | Gln | Thr | Glu | Glu 155 | Pro | His | Leu | | | | | | | | |
| | (2) | I | NFOF | TAMS (i) | S (| EQUE A) | ENCE LEN | CHI GTH | | TER | ISTI base | e pa | irs | | | |
| 40 | | | | (xi | (| B) C) D) SEQU | STR TOP | OLO | nuc EDNE GY: ESCR | SS: li | s: nea: | ingl r | | O NC |):40: | - |
| 45 | | | | | | Leu | | | | | Leu | | | | GCC Ala 15 | 45 |

| | GGG CT | | Leu P | | | | | | | | | | | 90 |
|-------------|------------------|----------------|---------------|-----------------------------------|---------------------------|------------------|-----------------------------|----------------------------|---------------------|-------------|------------|------------|------------------|------|
| 5 | GCT GG Ala Gl | | Gly S | | | | | | | | | | | 135 |
| 10 | GTG TG Val Tr | | Arg S | | | | | | | | | | | 180 |
| 15 | GTC GT Val Va | | Glu T | | | | | | | | | | | 225 |
| | TCC TG Ser Cy | T GTC s Val | Ser L | TG CTG eu Leu 80 | CGC Arg | TGC Cys | ACC Thr | GGC Gly 85 | TGC Cys | TGC Cys | G1y GGC | GAT Asp | GAG Glu 90 | 270 |
| 20 | AAT CT Asn Le | | Cys V | | | | | | | | | | | 315 |
| 25 | CTC CT Leu Le | | Ile A | | | | | | | | | | | 360 |
| | ACG TT Thr Ph | | Gln H | | | | | | | | | | | 405, |
| 30 | AAG AT Lys Me | | Pro G | | | | | | | | | | | 450 |
| 35 | AGG AG Arg Ar | | | AG | | | | | | | | | | 465 |
| 40 | (2) | INFO | (i) | SEQUI (A) (B) (C) (D) | ENCE LEN TYP STR | CHI GTH E: | ARAC : 1 amin EDNE | TER: 54 a o a SS: | ISTI amin cid | o a .ngl | | 6 | | |
| 45 . | Met Pr | o Val | (xi) Met A | SEQU rg Leu 5 | | | | • | | | | | | |
| 50 | Gly Le | u Ala | Leu P | ro Ala 20 | Val | Pro | Pro | Gln 25 | Gln | Trp | Ala | Leu | Ser 30 | |

Ala Gly Asn Gly Ser Ser Glu Val Glu Val Val Pro Phe Gln Glu 35 Val Trp Gly Arg Ser Tyr Cys Arg Ala Leu Glu Arg Leu Val Asp Val Val Ser Glu Tyr Pro Ser Glu Val Glu His Met Phe Ser Pro 70 65 10 Ser Cys Val Ser Leu Leu Arg Cys Thr Gly Cys Cys Gly Asp Glu Asn Leu His Cys Val Pro Val Glu Thr Ala Asn Val Thr Met Gln 100 95 15 Leu Leu Lys Ile Arg Ser Gly Asp Arg Pro Ser Tyr Val Glu Leu 115 Thr Phe Ser Gln His Val Arg Cys Glu Cys Arg Pro Leu Arg Glu 20 Lys Met Lys Pro Glu Arg Arg Pro Lys Gly Arg Gly Lys Arg Arg Arg Glu Lys

25

Claims

- A purified and isolated DNA sequence encoding the C subunit of vascular endothelial cell growth factor.
 - 2. A purified and isolated vascular endothelial cell growth factor C subunit DNA sequence comprising:

```
ATG CCG GTC ATG AGG CTG TTC CCT TGC TTC CTG CAG CTC CTG
35
       GOO GGG CTG GOG CTG COT GCT GTG COO COO CAG CAG TGG GCC
       TTG TCT GCT GGG AAC GGC TCG TCA GAG GTG GAA GTG GTA CCC
       TTC CAG GAA GTG TGG GGC CGC AGC TAC TGC CGG GCG CTG GAG
40
       AGG CTG GTG GAC GTC GTG TOC GAG TAC COC AGC GAG GTG GAG
       CAC ATG TTC AGC CCA TCC TGT GTC TCC CTG CTG CGC TGC ACC
       GGC TGC TGC GGC GAT GAG AAT CTG CAC TGT GTG CCG GTG GAG
       ACG GCC AAT GTC ACC ATG CAG CTC CTA AAG ATC CGT TCT GGG
       GAC CGG CCC TCC TAC GTG GAG CTG ACG TTC TCT CAG CAC GTT
       CGC TGC GAA TGC CGG CCT CTG CGG GAG AAG ATG AAG CCG GAA
       AGG AGG AGA COC AAG GGC AGG GGG AAG AGG AGG AGA GAG AAG
50
       TAG.
             SEQ ID NO:40
```

- Vascular endothelial cell growth growth factor AC DNA comprising an A subunit DNA sequence and a C subunit DNA sequence.
- Vascular endothelial cell growth growth factor BC DNA comprising a B subunit DNA sequence and a C subunit DNA sequence.

A purified and isolated vascular endothelial cell growth factor C subunit DNA sequence comprising:
 ATG COG GTC ATG AGG CTG TTC CCT TGC TTC CTG CAG CTC CTG

```
5
      GOO GOG CTG GOT GOT GOT GTG GOO GOO CAG GAG TGG GOO
       TIG ICI GCI GGG AAC GGC ICG ICA GAG GIG GAA GIG GIA CCC
       TTC CAG GAA GTG TGG GGC OGC AGC TAC TGC OGG GOG CTG GAG
      AGG CTG GTG GAC GTC GTG TOC GAG TAC COC AGC GAG GTG GAG
10
      CAC ATG TTC AGC CCA TOC TGT GTC TCC CTG CTG CGC TGC ACC
       GGC TGC TGC GGC GAT GAG AAT CTG CAC TGT GTG CCG GTG GAG
       ACG GCC AAT GTC ACC ATG CAG CTC CTA AAG ATC CGT TCT GGG
15
       GAC OGG COC TOC TAC GIG GAG CIG AOG TIC TCI CAG CAC GIT
       CGC TGC GAA TGC CGG CCT CTG CGG GAG AAG ATG AAG CCG GAA
       AGG AGG AGA COC AAG GGC AGG GGG AAG AGG AGG AGA GAG AAG
       CAG AGA COC ACA GAC TGC CAC CTG TGC GGC GAT GCT GTT CCC
20
       CGG AGG TAA.
                     SEQ ID NOS:29 & 40
```

- 6. Vascular endothelial cell growth factor AC DNA comprising an A subunit DNA sequence selected form the group consisting of: a DNA sequence encoding an 189 amino acid form, a DNA sequence encoding an 165 amino acid form and a DNA sequence encoding a 121 amino acid form, with said A subunit DNA operably attached to a C subunit DNA sequence.
- 7. Vascular endothelial cell growth factor BC DNA comprising a B subunit DNA sequence selected form the group consisting of a DNA sequence encoding a 135 amino acid form; and a DNA sequence encoding a 115 amino acid form, with said B subunit DNA sequence operably attached to a C subunit DNA sequence.
 - 8. Homodimeric vascular endothelial growth factor DNA comprising C subunit DNA sequences.
- 9. A vector containing the DNA sequence of any one of claims 3 to 8.
 - A host cell transformed by the vector of claim 9 containing the DNA sequence encoding vascular endothelial cell growth factor.
- 11. A process for the preparation of vascular endothelial cell growth factor comprising culturing the transformed host cell of claim 10 under conditions suitable for the expression of vascular endothelial cell growth factor and recovering vascular endothelial cell growth factor.
 - 12. Vascular endothelial growth factor made by the process of claim 11.
- 45 13. Vascular endothelial cell growth factor AC comprising an A subunit amino acid sequence and a C subunit amino acid sequence.
 - 14. Vascular endothelial cell growth factor BC comprising a B subunit amino acid sequence and a C subunit amino acid sequence.
- 15. Vascular endothelial cell growth factor CC comprising a C subunit amino acid sequence and a C subunit amino acid sequence.
 - A purified and isolated vascular endothelial cell growth factor C subunit amino acid sequence comprising:

Met Pro Val Met Arg Ieu Phe Pro Cys Phe Ieu Gln Ieu Ieu Ala Gly Ieu Ala Ieu Pro Ala Val Pro Pro Gln Gln Trp Ala Ieu Ser Ala Gly Asn Gly Ser Ser Glu Val Glu Val Val Pro Phe Gln Glu Val Trp Gly Arg Ser Tyr Cys Arg Ala Ieu Glu Arg Ieu Val Asp Val Val Ser Glu Tyr Pro Ser Glu Val Glu His Met Phe Ser Pro Ser Cys Val Ser Ieu Ieu Arg Cys Thr Gly Cys Cys Gly Asp Glu Asn Ieu His Cys Val Pro Val Glu Thr Ala Asn Val Thr Met Gln Ieu Ieu Iys Ile Arg Ser Gly Asp Arg Pro Ser Tyr Val Glu Ieu Thr Phe Ser Gln His Val Arg Cys Glu Cys Arg Pro Ieu Arg Glu Iys Met Iys Pro Glu Arg Arg Arg Arg Pro Iys Gly Arg Gly Iys Arg Arg Arg Glu Iys. SEQ ID NO: 41

20 17. A purified and isolated vascular endothelial cell growth factor C subunit amino acid sequence comprising:

Met Pro Val Met Arg Leu Phe Pro Cys Phe Leu Gln Leu Leu Ala Gly Leu Ala Leu Pro Ala Val Pro Pro Gln Gln Trp Ala 25 Leu Ser Ala Gly Asn Gly Ser Ser Glu Val Glu Val Val Pro Phe Gln Glu Val Trp Gly Arg Ser Tyr Cys Arg Ala Leu Glu Arg Leu Val Asp Val Val Ser Glu Tyr Pro Ser Glu Val Glu His Met Phe Ser Pro Ser Cys Val Ser Leu Leu Arg Cys Thr 30 Gly Cys Cys Gly Asp Glu Asn Leu His Cys Val Pro Val Glu Thr Ala Asn Val Thr Met Gln Leu Leu Lys Ile Arg Ser Gly Asp Arg Pro Ser Tyr Val Glu Leu Thr Phe Ser Gln His Val 35 Arg Cys Glu Cys Arg Pro Leu Arg Glu Lys Met Lys Pro Glu Arg Arg Arg Pro Lys Gly Arg Gly Lys Arg Arg Arg Glu Lys Gln Arg Pro Thr Asp Cys His Leu Cys Gly Asp Ala Val Pro 40 Arg Arg. SEQ ID NOS: 29 & 40

- 18. A tissue repairing pharmaceutical composition comprising a pharmaceutical carrier and an effective tissue repairing amount of the purified vascular endothelial growth factor of any one of claims 13 to 15,
- 19. The use of the vascular endothelial cell growth factor of any one of claims 13 to 15 for the manufacture of a medicament for promoting tissue repair.
- 20. The use of the vascular endothelial cell growth factor of any one of claims 13 to 15 for the manufacture of a medicament for stimulating vascular endothelial cell growth.

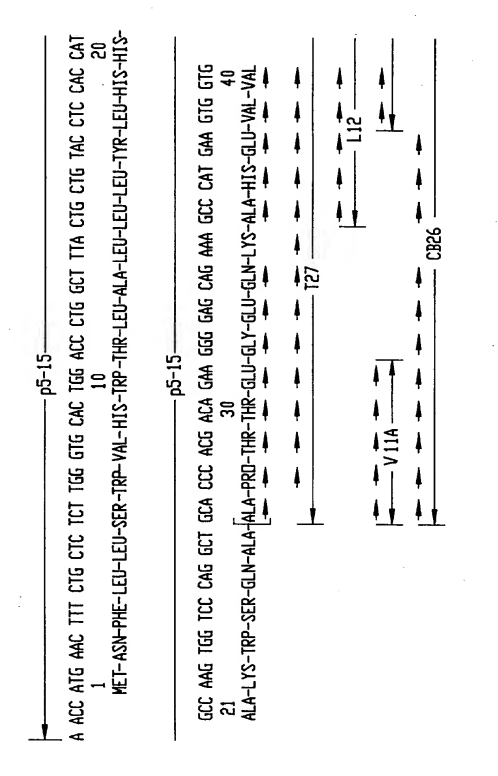
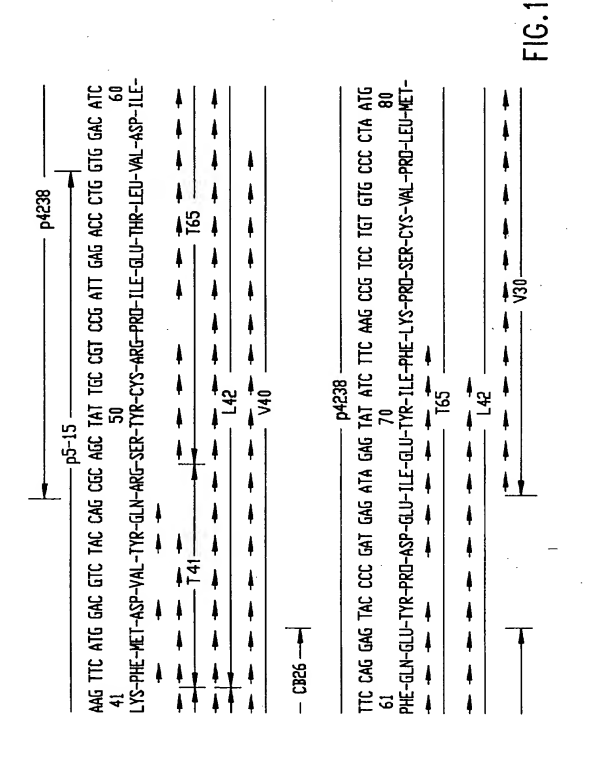
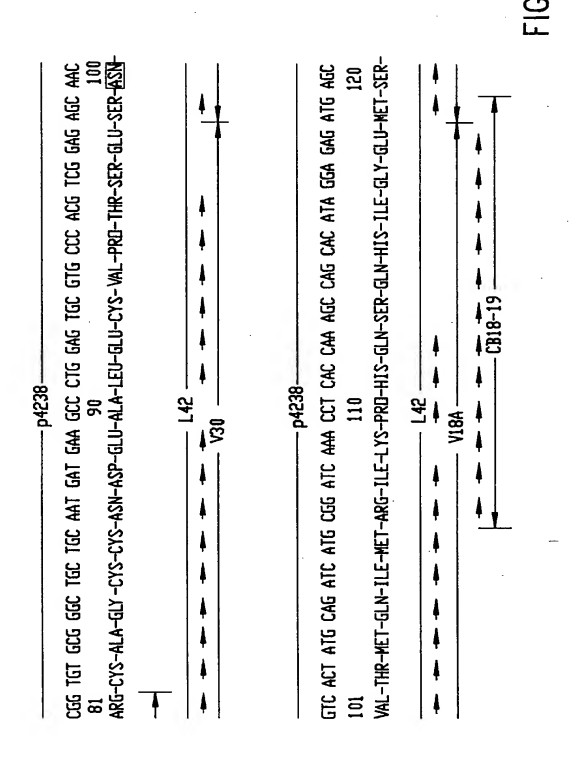
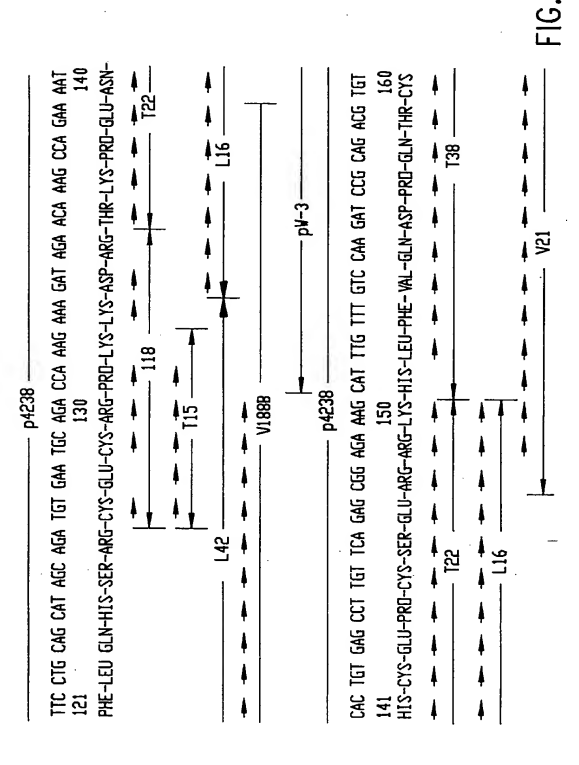
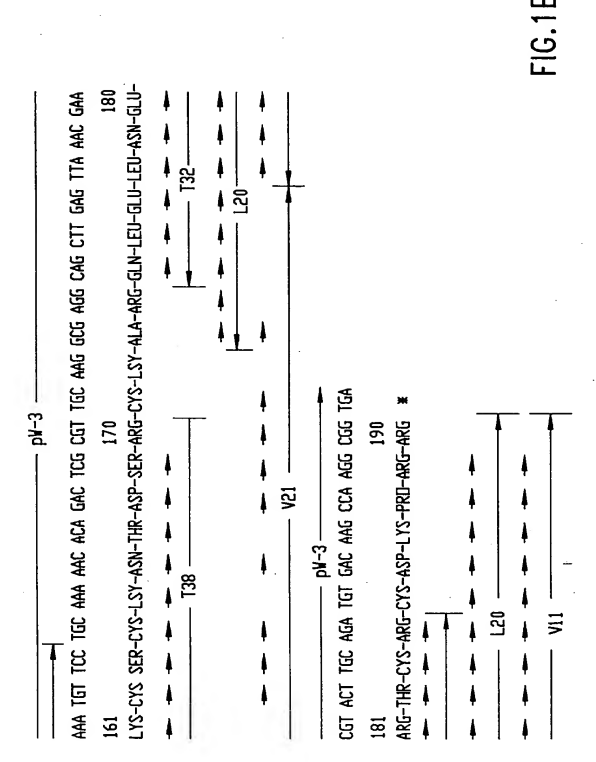


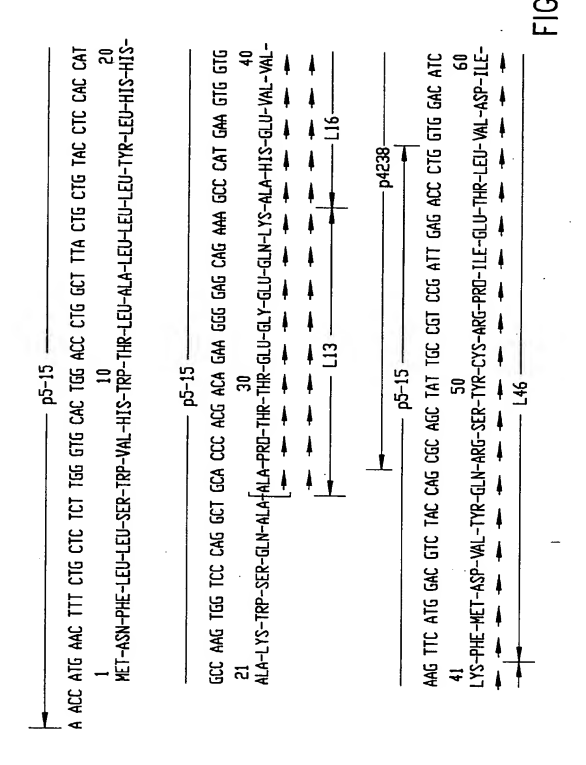
FIG 1A



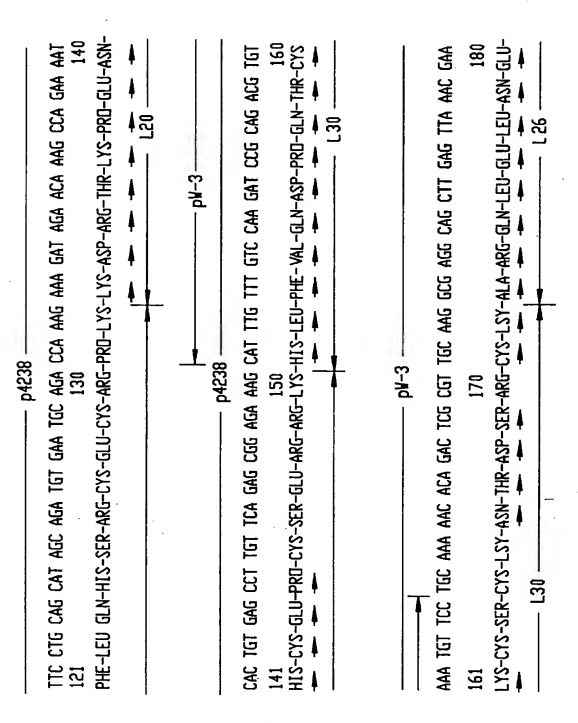


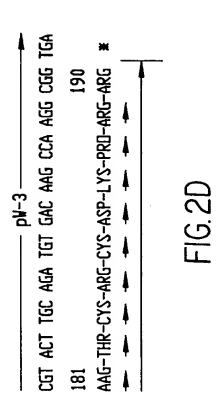


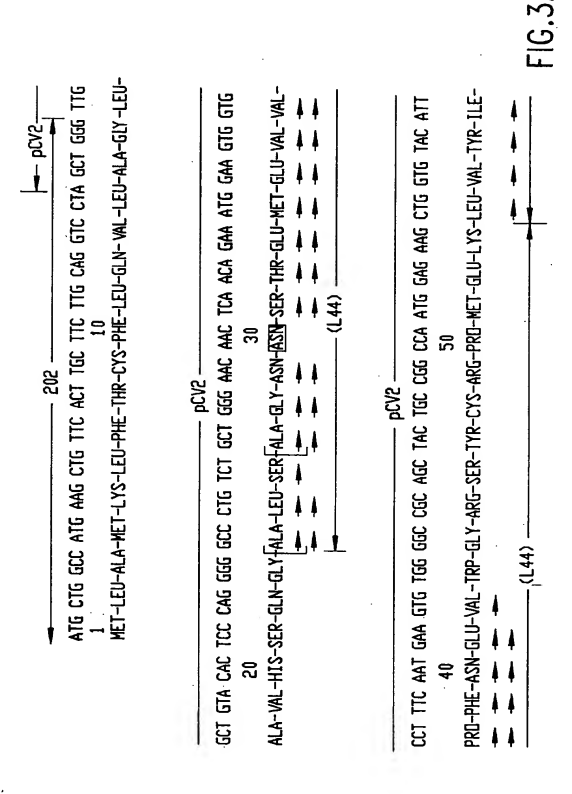


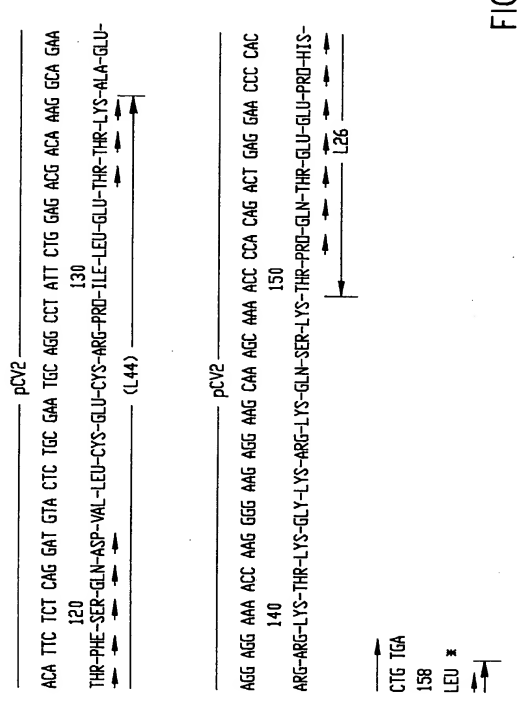


F1G.2C









| ACCA | ATG Met | AAC Asn | TTT Phe | CIG | CIC Leu 5 | TCT Ser | TGG | GIG Val | CAC His | TGG Trp 10 | Thr | CIG Leu | GCT Ala | TTA | CTG Leu 15 | 49 |
|------|--------------|------------|------------|------------|-------------------|------------|------------|------------|------------|-------------------|------------|------------|------------|------------|-------------------|---------|
| | CTG Leu | TAC Tyr | CTC Leu | CAC His | CAT His 20 | GCC Ala | AAG Lys | TGG | TCC Ser | CAG Gln 25 | GCT Ala | GCA Ala | occ Pro | ACG Thr | ACA Thr 30 | 94 |
| | GAA Glu | GGG Gly | GAG Glu | CAG Gln | AAA Lys 35 | GCC Ala | CAT His | GAA Glu | GTG Val | GTG Val 40 | AAG Lys | TTC Phe | ATG Met | GAC Asp | GTC Val 45 | 139 |
| | TAC Tyr | CAG Gln | CGC Arg | AGC Ser | TAT Tyr 50 | TGC Cys | OGT Arg | CCG Pro | ATT Ile | GAG Glu 55 | ACC Thr | CIG Leu | GTG Val | GAC Asp | ATC Ile 60 | 184 |
| | TTC Phe | CAG Gln | GAG Glu | TAC Tyr | CCC Pro 65 | GAT Asp | GAG Glu | ATA Ile | GAG Glu | TAT Tyr 70 | ATC Ile | TTC Phe | AAG Lys | CCG Pro | TCC Ser 75 | 229 |
| | TGT Cys | GTG Val | ccc Pro | CTA Leu | ATG Met 80 | CGG Arg | TGT Cys | GCG Ala | GC Gly | TGC Cys 85 | TGC Cys | AAT Asn | GAT QzA | GAA Glu | GCC Ala 90 | 274 |
| | CTG Leu | GAG Glu | TGC Cys | GTG Val | CCC Pro 95 | ACG Thr | TCG Ser | GAG Glu | AGC Ser | AAC Asn 100 | GTC Val | ACT Thr | ATG Met | CAG Gln | ATC Ile 105 | |
| | ATG Met | CGG Arg | ATC Ile | Lys | CCT Pro 110 | CAC His | CAA Gln | AGC Ser | CAG Gln | CAC His 115 | ATA Ile | GGA Gly | GAG Glu | Met | AGC Ser 120 | 364 |
| | TTC Phe | CIG Leu | CAG Gln | His | AGC Ser 125 | AGA Arg | TGT Cys | GAA Glu | TGC Cys | AGA Arg 130 | CCA Pro | AAG Lys | AAA Lys | Asp | AGA Arg 135 | 409 |
| | ACA . Thr | AAG Lys | CCA Pro | Glu | AAA Lys 140 | TGT Cys | GAC Asp | AAG Lys | CCA Pro | AGG Arg 145 | CCG Arg | TGA | | | | 445 |

FIG. 4

| AACC | ATG Met | AAC Asn | TTT Phe | CIG Leu | CTC Leu 5 | TCT Ser | TCG Trp | GIG Val | CAC His | TGG Trp 10 | ACC Thr | CIG Leu | OCT Ala | TTA Leu | CTG Leu 15 | 49 |
|------|------------|------------|------------|------------|-------------------|------------|------------|------------|------------|-------------------|------------|------------|------------|------------|-------------------|-----|
| | CTG Leu | TAC Tyr | CTC | CAC His | CAT His 20 | GCC Ala | AAG Lys | TGG Trp | TCC Ser | CAG Gln 25 | GCT Ala | GCA Ala | œ Pro | ACG Thr | ACA Thr 30 | 94 |
| | GAA Glu | GGG Gly | GAG Glu | CAG Gln | AAA Lys 35 | GCC Ala | CAT His | GAA Glu | GIG Val | GTG Val 40 | AAG Lys | TTC Phe | ATG Met | GAC Asp | GTC Val 45 | 139 |
| | TAC Tyr | CAG Gln | CCC Arg | AGC Ser | TAT Tyr 50 | TGC Cys | OGT Arg | ccc Pro | ATT Ile | GAG Glu 55 | ACC Thr | CTG Leu | GTG Val | GAC Asp | ATC Ile 60 | 184 |
| | TTC Phe | CAG Gln | GAG Glu | TAC Tyr | CCC Pro 65 | GAT Asp | GAG Glu | ATA Ile | GAG Glu | TAT Tyr 70 | ATC Ile | TTC Phe | AAG Lys | CCG Pro | TCC Ser 75 | 229 |
| | TGT Cys | GTG Val | ccc Pro | CTA Leu | ATG Met 80 | CGG Arg | TGT Cys | GOG Ala | GGC Gly | TGC Cys 85 | TGC Cys | AAT Asn | GAT Asp | GAA Glu | CCC Ala 90 | 274 |
| | CTG Leu | GAG Glu | TGC Cys | GIG Val | CC Pro 95 | ACG Thr | TCG Ser | GAG Glu | AGC Ser | AAC Asn 100 | GTC Val | ACT Thr | ATG Met | Gln | ATC Ile 105 | 319 |
| | ATG Met | CGG Arg | ATC Ile | AAA Lys | CCT Pro 110 | CAC His | CAA Gln | AGC Ser | CAG Gln | CAC His 115 | ATA Ile | GGA Gly | GAG Glu | Met | AGC Ser 120 | 364 |
| | TTC Phe | CTG Leu | CAG Gln | His | AGC Ser 125 | AGA Arg | TGT Cys | GAA Glu | TGC Cys | AGA Arg 130 | CCA Pro | AAG Lys | AAA Lys | Asp | AGA Arg 135 | 409 |
| | ACA Thr | AAG Lys | CCA Pro | Glu | AAT Asn 140 | CAC His | TGT Cys | GAG Glu | CCT Pro | TGT Cys 145 | TCA Ser | GAG Glu | CCG Arg | Arg | AAG Lys 150 | 454 |
| ٠ | CAT His | TTG Leu | TTT Phe | Val | CAA Gln 155 | GAT Asp | CCG Pro | CAG Gln | ACG Thr | TGT Cys 160 | AAA Lys | TGT Cys | TCC Ser | Cys | AAA Lys 165 | 499 |
| | AAC Asn | ACA Thr | GAC Asp | Ser | CGT Arg 170 | TGC Cys | AAG Lys | CCG Ala | AGG Arg | CAG Gln 175 | CTT Leu | GAG Glu | TTA Leu | Asn | GAA Glu 180 | 544 |
| | OGT Arg | | | Arg | | | | | | | TGA | ł | -1(- | : 5 | | 577 |

| AACC | ATG Met | AAC Asn | TTT Phe | CIG Leu | CTC Leu 5 | TCT Ser | TGG Trp | GTG Val | CAC His | TGG Trp 10 | ACC Thr | CTG Leu | GCT Ala | TTA Leu | CTG Leu 15 | 49 |
|------|------------|------------|------------|------------|-------------------|------------|------------|------------|------------|-------------------|------------|------------|------------|------------|-------------------|---------|
| | CTG Leu | TAC Tyr | CTC Leu | CAC His | CAT His 20 | GCC Ala | AAG Lys | TGG Trp | TCC Ser | CAG Gln 25 | GCT Ala | GCA Ala | ccc Pro | ACG Thr | ACA Thr 30 | 94 |
| | GAA Glu | GJy GGG | GAG Glu | CAG Gln | AAA Lys 35 | CCC Ala | CAT His | GAA Glu | GTG Val | GIG Val 40 | AAG Lys | TTC Phe | ATG Met | GAC Asp | GTC Val 45 | 139 |
| | TAC Tyr | CAG Gln | CGC Arg | AGC Ser | TAT Tyr 50 | TGC Cys | CGT Arg | CCG Pro | ATT Ile | GAG Glu 55 | ACC Thr | CTG Leu | GTG Val | GAC Asp | ATC Ile 60 | 184 |
| | TTC Phe | CAG Gln | GAG Glu | TAC Tyr | CCC Pro 65 | GAT Asp | GAG Glu | ATA Ile | GAG Glu | TAT Tyr 70 | ATC Ile | TTC Phe | AAG Lys | CCG Pro | TCC Ser 75 | |
| | TGT Cys | GTG Val | CCC Pro | CTA Leu | ATG Met 80 | CGG Arg | TGT Cys | GCG Ala | GGC Gly | TGC Cys 85 | TGC Cys | AAT Asn | GAT Asp | GAA Glu | GCC Ala 90 | 274 |
| | CTG Leu | GAG Glu | TGC Cys | GIG Val | ccc Pro 95 | ACG Thr | TCG Ser | GAG Glu | AGC Ser | AAC Asn 100 | GTC Val | ACT Thr | ATG Met | Gln | ATC Ile 105 | 319 |
| | ATG Met | CGG Arg | ATC Ile | Lys | CCT Pro 110 | CAC His | CAA Gln | AGC Ser | CAG Gln | CAC His 115 | ATA Ile | GGA Gly | GAG Glu | Met | AGC Ser 120 | 364 |
| | TTC Phe | CTG Leu | CAG Gln | His | AGC Ser 125 | AGA Arg | TGT Cys | GAA Glu | TGC Cys | AGA Arg 130 | CCA Pro | AAG Lys | AAA Lys | Asp | AGA Arg 135 | 409 |
| | ACA Thr | AAG Lys | CCA Pro | Glu | AAA Lys 140 | AAA Lys | TCA Ser | GIT Val | CGA Arg | GGA Gly 145 | AAG Lys | GGA Gly | AAG Lys | Gly | CAA Gln 150 | 454 |
| | AAA Lys | CGA Arg | AAG Lys | Arg | AAG Lys 155 | AAA Lys | TCC Ser | OGG Arg | TTT Phe | AAA Lys 160 | TCC Ser | TGG Trp | AGC Ser | Val | CAC His | 499 |

FIG. 6A

| GAG Glu | | | | | | | | GAT Asp 180 | 544 |
|------------|--|-----|--|--|--|--|-----|-------------------|-----|
| CAG Gln | | | | | | | | TGC Cys 195 | 589 |
| GCG Ala | | | | | | | Cys | GAC Asp 210 | 634 |
| CCA Pro | | TGA | | | | | | | 649 |

FIG.6B

| ATG Met | CTG Leu | GCC Ala | ATG Met | AAG Lys 5 | CTG Leu | TTC Phe | ACT Thr | TGC Cys | TTC Phe 10 | TTG Leu | CAG Gln | GTC Val | CTA Leu | GCT Ala 15 | 45 |
|------------|------------|------------|------------|-------------------|------------|------------|------------|------------|-------------------|------------|------------|------------|------------|-------------------|-----|
| GGG Gly | TTG Leu | GCT Ala | GIG Val | CAC His 20 | TCC Ser | CAG Gln | GGG Gly | GCC Ala | CTG Leu 25 | TCT Ser | GCT Ala | GGG Gly | AAC Asn | AAC Asn 30 | 90 |
| TCA Ser | ACA Thr | GAA Glu | ATG Met | GAA Glu 35 | GTG Val | GTG Val | CCT Pro | TTC Phe | AAT Asn 40 | GAA Glu | GIG Val | TGG Trp | GC Gly | CCC Arg 45 | 135 |
| AGC Ser | TAC Tyr | TGC Cys | CGG Arg | CCA Pro 50 | ATG Met | GAG Glu | AAG Lys | CTG Leu | GTG Val 55 | TAC Tyr | ATT Ile | GCA Ala | GAT Asp | GAA Glu 60 | 180 |
| CAC His | CCT Pro | AAT Asn | GAA Glu | GTG Val 65 | TCT Ser | CAT His | ATA Ile | TTC Phe | AGT Ser 70 | CCG Pro | TCA Ser | TGT Cys | GTC Val | CTT Leu 75 | 225 |
| CIG Leu | AGT Ser | CGC Arg | TGT Cys | AGT Ser 80 | GGC Gly | TGC Cys | TGT Cys | Gly | GAC Asp 85 | GAG Glu | GT Gly | CTG Leu | CAC His | TGT Cys 90 | 270 |
| | | | | ACA Thr 95 | | | | | | | | | Lys | | 315 |
| | | | | GAT Asp 110 | | | | | | | | | Phe | | 360 |
| CAG Gln | GAT Asp | GTA Val | Leu | TGC Cys 125 | GAA Glu | TGC Cys | AGG Arg | CCT Pro | ATT Ile 130 | CIG Leu | GAG Glu | ACG Thr | Thr | AAG Lys 135 | 405 |
| | GAA | | TAA | | | | | | | | | | | • | 417 |

FIG.7

| ATG Met | CIG Leu | GCC Ala | ATG Met | AAG Lys | CTG Leu | TTC Phe | ACT Thr | TGC Cys | Phe | MG Leu | CAG Gln | GTC Val | CTA Leu | Ala | 45 |
|------------|------------|------------|------------|-------------------|------------|------------|------------|------------|-------------------|------------|------------|-------------|------------|-------------------|-----|
| GGG Gly | TTG Leu | GCT Ala | GTG Val | CAC His 20 | TCC Ser | CAG Gln | GGG Gly | GOC Ala | CTG Leu 25 | TCT Ser | GCT Ala | GJ y GGG | AAC Asn | AAC Asn 30 | 90 |
| TCA Ser | ACA Thr | GAA Glu | ATG Met | GAA Glu 35 | GTG Val | GIG Val | CCT Pro | TTC Phe | AAT Asn 40 | GAA Glu | GTG Val | TGG Trp | GGC Gly | OGC Arg 45 | 135 |
| AGC Ser | TAC Tyr | TGC Cys | CGG Arg | CCA Pro 50 | ATG Met | GAG Glu | AAG Lys | CTG Leu | GTG Val 55 | TAC Tyr | ATT Ile | GCA Ala | GAT Asp | GAA Glu 60 | 180 |
| CAC His | CCT Pro | AAT Asn | GAA Glu | GTG Val 65 | TCT Ser | CAT His | ATA Ile | TTC Phe | AGT Ser 70 | CCG Pro | TCA Ser | TGT Cys | GTC Val | CTT Leu 75 | 225 |
| CTG Leu | AGT Ser | CGC Arg | TGT Cys | AGT Ser 80 | GGC Gly | TGC Cys | TGT Cys | GGT Gly | GAC Asp 85 | GAG Glu | GGT Gly | CTG Leu | CAC His | TGT Cys 90 | 270 |
| GIG Val | GOG Ala | CTA Leu | AAG Lys | ACA Thr 95 | GCC Ala | AAC Asn | ATC Ile | ACT Thr | ATG Met 100 | CAG Gln | ATC Ile | TTA Leu | Lys | ATT Ile 105 | 315 |
| cc Pro | cc Pro | AAT Asn | CGG Arg | GAT Asp 110 | CCA Pro | CAT His | TCC Ser | TAC Tyr | GTG Val 115 | GAG Glu | ATG Met | ACA Thr | Phe | TCT Ser 120 | 360 |
| CAG Gln | GAT Asp | GTA Val | Leu | TGC Cys 125 | GAA Glu | TGC Cys | AGG Arg | CCT Pro | ATT Ile 130 | CTG Leu | GAG Glu | AOG Thr | Thr | AAG Lys 135 | 405 |
| GCA Ala | GAA Glu | AGG Arg | Arg | AAA Lys 140 | ACC Thr | AAG Lys | œ Gly | aag Lys | AGG Arg 145 | AAG Lys | CAA Gln | AGC Ser | Lys | ACC Thr 150 | 450 |
| CCA Pro | CAG Gln | ACT Thr | Glu | GAA Glu 155 | CCC Pro | CAC His | CTG Leu | TGA | | | | | | | 477 |

FIG. 8

| Pro | | | | | | | | | 45 |
|------------|-----|-----|--|--|-----|--|--|-----|-----|
| CTG Leu | | | | | CAG | | | TCT | 90 |
| GGG Gly | | | | | | | | | 135 |
| TGG Trp | | | | | | | | | 180 |
| GTG Val | | | | | | | | | 225 |
| TGT Cys | | | | | | | | | 270 |
| CIG Leu | | | | | | | | | 315 |
| CTA Leu | Ile | | | | | | | | 360 |
| TTC Phe | Gln | | | | | | | | 405 |
| ATG Met | Pro | | | | | | | | 450 |
| AGA Ara | | TAG | | | | | | | 465 |

FIG. 9



EUROPEAN SEARCH REPORT

Application Number

EP 92 30 2750

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